Effects of solar irradiation on thermally driven CO₂ methanation using Ni/CeO₂–based catalyst

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Abstract

Utilization of the renewable energy sources is one of the main challenges in the state-of-the-art technologies for CO₂ recycling. Here we have taken advantage of the solar light harvesting in the thermocatalytic approach to carbon dioxide methanation. The large-surface-area Ni/CeO₂ catalyst produced by a scalable low-cost method was characterized and tested in the dark and under solar light irradiation conditions. Light-assisted CO₂ conversion experiments as well as in-situ DRIFT spectrometry, performed at different illumination intensities, have revealed a dual effect of the incident photons on the catalytic properties of the two-component Ni/CeO₂ catalyst. On the one hand, absorbed photons induce a localized surface plasmon resonance in the Ni nanoparticles followed by dissipation of the heat to the oxide matrix. On the other hand, the illumination activates the photocatalytic properties of the CeO₂ support, which leads to an increase in the concentration of the intermediates being precursor for methane production. Analysis of the methane production at different temperatures and illumination conditions has shown that the methanation reaction in our case is controlled by a photothermally-activated process. The used approach has allowed us to increase the reaction rate up to 2.4 times and consequently decrease the power consumption by 20 % under solar illumination, thus replacing the conventional thermal activation of the reaction with a green energy source.

1. Introduction

Conversion of carbon dioxide to renewable hydrocarbon fuels, such as methane, is a big challenge nowadays due to the high thermodynamic stability of the CO₂ molecule and, therefore, major difficulties in its reduction [1–3]. The standard thermocatalytic approach for CO₂ methanation requires relatively high temperatures (150–500 °C) [4], which gives rise to a number of alternative approaches to replace or assist thermochemical catalysis [5–8]. Plasma-assisted methanation of CO₂ was reported, being efficient even at room temperature [9,10] with promising performances. Another important field of research is related to the utilization of solar light as an additional green energy source for CO₂ hydrogenation to methane [11–15]. Except for the standard photocatalytic approach, based on the metal-decorated metal oxides, where the metal oxide support is used as the main photoabsorber and metallic nanoparticles act only as active catalytic sites [16,17], a rapidly emerging field in catalysis is taking advantage of the strong light extinction of the metallic nanoparticles themselves, owning to the effect called localized surface plasmon resonance (LSPR) [18,19]. The LSPR-driven catalysis is mostly making use of noble metals (Au and Ag) due to their strong plasmonic response in the visible and UV part of the spectrum [20–24]. However, other abundant and non-expensive metals, in particular some of the VIII group transition metals, such as Ni have also proven to exhibit plasmonic features [11,25–29], although less pronounced due to the higher damping. Moreover, nickel nanoparticles in combination with metal oxides presenting high oxygen mobility, such as CeO₂, have been proposed as a candidate catalyst for the efficient thermal conversion of CO₂ to methane [30].

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Excited LSPR is followed by a strong near-field enhancement, which is then damped through one of the possible mechanisms: i) radiative decay through re-emitting of the photons; ii) Landau damping, resulting in a non-equilibrium charge distribution (so-called “hot electrons” or “hot holes”, which then can be transferred to the metal-oxide support or directly participate in the catalytic reaction); or iii) direct excitation of electron-hole pairs in the semiconductor through plasmon-induced resonant energy transfer (PIRET). Another effect that has been recently introduced, consists in direct metal-to-semiconductor charge transfer [31,32]. All of the damping mechanisms stated above occur in the timescale of 1–100 fs and, eventually, lead to recombination of the created non-equilibrium charge carriers (0.1–100 ps) and result in strong local heating (0.1–10 ns) [33]. The plasmon damping has found a lot of applications, and many works are dedicated to exploiting of the “direct” improvement of catalytic properties by plasmonic hot electrons [20,25,34–37].

However, few works have emphasized the importance of the catalytic performance enhancement of the Ni-based catalysts due to plasmon-induced self-heating of nickel nanoparticles followed by a dissipation of heat to the oxide matrix. Due to the excellent photo-thermal properties of Ni nanoparticles [26,38–40], this effect in many cases can be considered as the main consequence of the plasmon excitation in Ni-based catalysts as compared with direct hot electrons interface transfer. In this work, we have used a wet impregnation of mesoporous silica SBA-15 template to prepare a high surface area Ni/CoO2 catalyst for CO2 methanation where a large size distribution of Ni nanoparticles allows us to perform highly selective and energetically efficient methanation of CO2 assisted by plasmonic heating of the catalysts with solar light. The use of in-situ characterization has revealed the mechanism of CO2 hydrogenation and the optimal conditions for the plasmon-assisted methanation reaction, which allows us to increase the productivity up to 2.4 times and reduce the energy consumption by 20 % compared to the conventional thermochemical approach. Furthermore, we have corroborated that under solar illumination conditions, in spite of the significant increase of the formate intermediates due to the mechanism of CO2 hydrogenation and the optimal conditions for the plasmon-assisted methanation reaction, which allows us to increase the productivity up to 2.4 times and reduce the energy consumption by 20 % compared to the conventional thermochemical approach.

2. Experimental

2.1. Catalyst synthesis

SBA-15 mesoporous silica templates were synthesized by the following process, based on the one described in [41]: 9 g of Pluronic® P123 (polyethylene glycol-block-polypropylene glycol-block-polyethylene glycol, average Mn ~5,800, Sigma-Aldrich) was mixed with 90 g of diluted HCl (50 wt.%) and 245 g of Milli-Q water and stirred at 38 °C to obtain polymeric micelles. 18.7 g of TEOS (tetraethyl orthosilicate, 98 %, Sigma-Aldrich) was added as a silica precursor. The reaction was held for another 24 h at 100 °C, after which the mixture was filtered and eventually calcined at 550 °C for 5 h.

The Ni/CoO2 catalyst was prepared by simultaneous impregnation of SBA-15 templates (1.05 g) with 6.5 mL of 0.6 M cerium nitrate hexahydrate (Sigma-Aldrich) and 3 mL of 0.6 M nickel (II) nitrate hexahydrate (Sigma-Aldrich). The Ni/Co ratio was chosen in such a way that the estimated Ni weight concentration in the final product would be 15 %. The reaction was held in two steps, to achieve a high degree of impregnation [41,42]: first with impregnation of the solutions to the mesoporous silica and annealing at 350 °C for the preliminary precursor decomposition, and second impregnation with half of the initial amounts, followed by annealing of the sample at 800 °C. The silica template was further dissolved in 2 M NaOH for 12 h at 70 °C with constant stirring. The obtained NiO/CoO2 powder was further reduced in hydrogen atmosphere (5% H2 and 95 % Ar) for 7 h at 450 °C to get the Ni/CoO2 catalyst (Scheme 1).

For comparison, samples without Ni were also prepared using the same template impregnation technique. 6.5 mL of 0.6 M cerium nitrate hexahydrate was added to 0.9 g of SBA-15, and the same two-step procedure was applied for calcination and second impregnation of silica template.

2.2. Physico-chemical characterization

XRD spectra were recorded on a Bruker type XRD D8 diffraction-meter (CuKα radiation, λ = 1.5418 Å, 40 kV, 40 mA) with a scanning range of 2θ from 20° to 80° and step size of 0.05° / 3 s. The average crystalline sizes of metallic nanoparticles were derived from the Scherrer’s equation \( D = \frac{K \lambda}{\beta \cos \theta} \), where \( \Theta \) is the Bragg angle, \( \lambda \) is the X-ray wavelength and \( \beta \) is the full width of the diffraction peak at half maximum (FWHM).

Scanning electron microscope (SEM, Zeiss Auriga 60) was used to study the morphology of the samples. The composition of the synthesized catalyst was studied by energy dispersive X-ray spectroscopy (EDX, Oxford Inca Energy).

N2-physisorption analysis was conducted in TriStar II 3020-Micrometrics equipment at liquid nitrogen temperature. Before the measurements, the samples were degassed under vacuum conditions at 90 °C for 1 h and then at 250 °C for 3 h. Brunauer-Emmett-Teller (BET) method was applied for calculation of the BET surface area for a relative pressure (P/P0) range of 0.1–0.3. The total pore volume and the average pore size were determined by applying Barrett-Joyner-Halenda (BJH) method to desorption branch of the isotherms at the value P/P0 = 0.95.

H2-TPR (hydrogen temperature programmed reduction) and CO-chemisorption were carried out using an automated chemisorption analyzer (Autochem HP-Micrometrics). In case of H2-TPR measurements 100 mg of powder sample was kept under constant 5% H2/Ar flow, while the temperature was increased from 25 °C to 800 °C with a heating ramp of 10 °C/min. The consumed H2 was measured using a thermal conductivity detector (TCD). The CO-chemisorption measurements were held at 35 °C under the flow of 10 %CO/He over the samples, which were pre-reduced at 450 °C under 5% H2/Ar flow for 7 h. Pulses of CO were periodically introduced until full saturation of the system. Active metal surface area and metal dispersion were calculated assuming the stoichiometric ratio CO/Ni equal to 1, Ni atomic weight to 58.71, its metal surface area and metal dispersion were calculated assuming the stoichiometric ratio CO/Ni equal to 1, Ni atomic weight to 58.71, its density to 8.9 g/cm3.

UV–vis light absorption spectra were measured using the Perkin Elmer Lambda 35 UV–vis spectrometer in the diffuse reflectance mode. Absorption of the samples was calculated according to the equation: \( A = 1 – R \), as no transmission was observed.

High-resolution transmission electron microscopy (HRTEM) together with scanning TEM (STEM) investigation was performed on a field emission gun FEI Tecnai F20 microscope. High angle annular dark-field (HAADF) STEM was combined with electron energy loss spectroscopy (EELS) in the Tecnai microscope by using a GATAN QUANTUM filter.

2.3. Functional characterization

A custom-made flow-type gas reactor (Scheme S1(b)) was created in order to allow simultaneous heating and illumination of the catalyst. 50 mg of the catalyst were spread uniformly over an aluminum support, whereas its temperature was constantly measured by a K-type thermocouple. An intimate contact was preserved throughout the reaction in order to measure directly the changes in the catalyst temperature.

The catalyst was preliminarily reduced in hydrogen atmosphere (5% H2 and 95 % Ar) for 7 h at 450 °C. The methanation reaction was held under a constant flow of 80 % H2 and 20 % CO2 gas mixture (99.999 %,
Linde) at a rate of 20 mL min\(^{-1}\). The outlet gas flow passed through a cold liquid-gas separator (5 °C) for water condensation, and was measured by a mass flow meter (MF, Bronkhorst). The gas products (CH\(_4\), CO, CO\(_2\) and H\(_2\)) were analyzed using a 490 microGC gas chromatograph (Agilent Technologies).

The methane yield was calculated according to the equation:

\[
Y_{CH_4}(\%) = \frac{X_{CO_2}(\%)}{100}\times S_{CH_4}(\%)
\]

where \(X_{CO_2}\) is the conversion of CO\(_2\) and \(S_{CH_4}\) is the selectivity to methane. \(X_{CO_2}\) and \(S_{CH_4}\) are the CO\(_2\) molar flows in the inlet and in the outlet, respectively, and \(S_{CH_4}\) is the CH\(_4\) molar flow in the outlet. The catalytic activity was then evaluated from the following equation:

\[
k = \frac{\Phi_{CH_4} \times Y_{CH_4}(\%)}{m_i \times 100%}
\]

The reported data were the average values, taken from five measurements for each temperature point, with the measurement error less than 3%. Carbon mass balance was closed within the accuracy of ± 7%.

As a source of illumination, we used a 300 W solar simulator (Xe lamp, AM1.5 G filter) with a variable illumination power. The power density was measured by a laser powermeter (Gentec-EO UNO) containing a calibrated reference silicon cell. Bandpass filters (FGL400S, FGS900S, ThorLabs) were introduced to the system to cut a part of the spectrum as shown in Scheme S1(b).

Provided that methanation is an exothermic reaction, the temperature was allowed to reach a stable value in all of the studied conditions before any illumination was introduced.

### 2.4. In-situ DRIFTS characterization

In-situ diffuse reflectance infrared Fourier transform spectroscopy (DRIFTS) was used in order to study the nature of intermediate species formed during the methanation reaction over the Ni/CeO\(_2\) catalyst under dark and light-assisted reaction conditions. The spectra were recorded using Bruker-Vertex70 spectrophotometer equipped with a high-temperature reaction control cell with two ZnSe windows, one quartz window and a catalytic bed. The catalyst was pre-reduced at 450 °C under 5% H\(_2\)/He flow of 50 mL/min for 1 h, then purged in He gas for 40 min to remove the residual H\(_2\) from the cell and finally cooled down to desired temperature. The background spectrum was recorded in He at room temperature. CO\(_2\) methanation reaction was performed in 20 mL/min of gas mixture (75 % of stoichiometric H\(_2\)/CO\(_2\) ratio = 4 and 25 % of He balance) at 100 °C, 150 °C, 200 °C, 225 °C and 250 °C. Each measurement was recorded after 30 min in isothermal conditions, and followed by purging in He for 30 min to remove the excess water molecules. Spectra were recorded from 4200 cm\(^{-1}\) to 1000 cm\(^{-1}\) with a step of 1 cm\(^{-1}\). Thermal analyzer Thermostat was used to control the product stream by their molecular weights (m/s ratios). The illumination (5 suns) during the DRIFTS measurement was applied through the quartz window from the solar simulator used during the functional characterization.

### 3. Results and discussion

#### 3.1. Physical characterization of the catalyst

SEM and TEM images (Fig. 1) show that the obtained catalyst consists of CeO\(_2\) nanofibers (with a diameter of ~10 nm) and well-distributed Ni nanoparticles with the size distribution approximately from 5 to 50 nm. The EDX analysis shows the presence of 12 % (w.t.) Ni, which is in a good proximity with the initial estimations. Some residual silicon was spotted, which supposedly comes from the non-dissolved SBA-15 template, but its amount was as low as 1% of the catalyst mass.

The ceria support, shown in Fig. 1(d), has a typical channel shape due to the SBA-15 template where each channel is 8 nm distant to the closest. This configuration of the support leads to its mesoporous properties and allows the small Ni clusters to penetrate inside the ceria structure.

HRTEM study has shown that the used synthesis procedure is suitable to create a ceria-supported catalyst with a presence of highly distributed small Ni nanoparticles (~5 nm) which have a strong interaction with the ceria support (Fig. 2). They are mostly located in the voids between the ceria rods. CO\(_2\) methanation over Ni/CeO\(_2\) is a complex reaction, requiring simultaneous H\(_2\) and CO\(_2\) activation. The former occurs on the metallic Ni sites and the latter on the basic sites of CeO\(_2\) [43]. The challenge of the catalyst design is that products of these two reaction steps (H\(^+\) and CO\(_2\)) are separated by a spatial constraint, which has to be minimized in order to achieve better catalytic performance. From this point of view, highly distributed Ni nanoclusters embedded in the mesoporous CeO\(_2\) (Fig. 1(d)) represent the optimal catalyst configuration for CO\(_2\) methanation. We assume that the high exposure of the boundary phase Ni/CeO\(_2\) to the gas is responsible for the significant CO\(_2\) conversion.

From the low-magnification images (Figs. 1(d); S1(a), Ni panel) we can also see the presence of bigger Ni clusters (~20–50 nm). They are located mostly on the surface of the catalyst and therefore play the major role in the solar light absorption and scattering. Relatively high distribution of the nanoclusters’ sizes is responsible for broadening of the plasmonic extinction peak and black coloring of the reduced powder.

The N\(_2\)-physiosorption analysis of the silica SBA-15 template, CeO\(_2\) and Ni/CeO\(_2\) replica has shown that the obtained powders possess a mesoporous structure with a high BET surface area. The surface areas and pore dimensions of pure CeO\(_2\) support and Ni/CeO\(_2\) catalyst used in this study are quite similar, which indicates that Ni impregnation has not influenced the mesoporous morphology of the ceria support. Moreover, the Ni/CeO\(_2\) catalyst exhibits a good dispersion of Ni nanoparticles with high metal surface area, which was derived from CO-chemisorption analysis (Table 1).

H\(_2\)-TPR analysis was used to study reduction properties of the obtained catalyst (Fig. 3). The Ni/CeO\(_2\) sample presents a strong peak at T = 328 °C and two smaller ones at T = 213 °C and T = 271 °C, attributed to the reduction of the weakly bound NiO species on the CeO\(_2\) surface [44]. The more intensive one is related to the reduction of smaller nanoparticles, whereas the low-temperature peaks are due to the large NiO particles on the CeO\(_2\) support. The peak at T = 442 °C was assigned to the reduction of strongly bound NiO [45], and the high-temperature process...
at $T_\text{=}697\,^\circ\text{C}$ was attributed to the $\text{Ce}^{4+}$ to $\text{Ce}^{3+}$ reduction in $\text{CeO}_2$ [46].

The $\text{H}_2$-TPR analysis confirmed that the reduction temperature of $450\,^\circ\text{C}$ (dashed line on Fig. 3), chosen for $\text{NiO}/\text{CeO}_2$ catalyst is sufficient to reduce all the types of $\text{NiO}$ clusters. EELS chemical composition maps prove that the Ni present in the reduced sample is mostly metallic (Fig. S1).

Fig. 4 shows XRD spectra of the as-prepared samples and reduced under $\text{H}_2$ atmosphere (5% $\text{H}_2$ in Ar, 7 h, $450\,^\circ\text{C}$). As we can see from the figure, $\text{NiO}$ phase, that was present in the as-prepared catalyst, disappears completely in favor of metallic Ni. The peaks attributed to $\text{CeO}_2$ are shifted with respect to the clear $\text{CeO}_2$ phase in case of the non-reduced samples, which indicates the incorporation of Ni atoms and nanoclusters to $\text{CeO}_2$, causing stress onto the lattice [47]. This shift can be clearly seen on the inset of Fig. 4, where the $\text{CeO}_2$ diffraction peak (56.34$\,^\circ$) present in both pure $\text{CeO}_2$ and reduced $\text{NiO}/\text{CeO}_2$ samples, is shifted to 56.41$\,^\circ$ in the case of non-reduced $\text{NiO}/\text{CeO}_2$. After reducing, all Ni is removed from the $\text{CeO}_2$ lattice, which results in shifting of the peaks back to the positions of the $\text{CeO}_2$ phase. It is also worth noticing, that the peak is broadened with respect to the pure $\text{CeO}_2$ support, which indicates the stress and the lattice parameter inhomogeneity due to the presence of the Ni atoms and nanoclusters inside of the $\text{CeO}_2$ rods. The crystallite sizes of Ni particles were estimated by Sherrer’s equation using the peak of 2$\Theta$ = 44.51$\,^\circ$. The average size was found to be 27.9 nm, which is in accordance with the nanoparticle dimensions extracted from the TEM data.

In order to analyze the possible strain influence of the Ni nanoparticles embedded in the $\text{CeO}_2$ mesoporous channels, we used the Geometrical Phase Analysis (GPA) routines in the Digital Micrograph software [48]. Our GPA analyses showed a relative compression of the $\text{CeO}_2$ lattice planes when they are close to a Ni nanoparticle (Fig. S2). The profile obtained along the dark blue arrow pointed out in the figure corroborates that the CeO$_2$ (111) planes suffer a 4–5 % compression close to the Ni nanoparticle, which is in line with the previous XRD analysis.

3.2. Catalytic performance of the catalyst in the CO$_2$ methanation reaction

We performed functional characterization of the synthetized catalyst in the tubular-type reactor in dark conditions (Scheme S1(a)). The measured catalytic activity at different reaction temperatures is shown on the Fig. 5(a). We found that CO$_2$ conversion reaches 80 % at 300 $^\circ$C, and the selectivity to methane was as high as 95 %.

Due to its high surface area the catalytic activity of Ni/CeO$_2$ catalyst used in this work is comparable to the activity of the other Ni/CeO$_2$ catalysts reported previously [49,50].

We have also carried out a stability test which has revealed a high stability of the catalyst under constant reaction conditions at $T_\text{=}270\,^\circ\text{C}$ (Fig. 5(b)).

3.3. Photothermal effects

UV–vis spectroscopy was used to measure the reflectance of bare CeO$_2$ and Ni/CeO$_2$ samples. The obtained data were transformed into absorption as no transmittance was observed. As it can be seen from the Fig. 6(a), bare cerium oxide powder does not absorb under IR and visible illumination, but starts absorbing strongly starting from 430 nm, which corresponds to the reported band gap of oxygen vacancy-enriched CeO$_2$ nanoparticles (2.87 eV) [51]. In contrast, samples with Ni nanoparticles exhibit an extraordinary high absorption over all of the measured illumination range. We attribute this significant rise in the visible and IR light absorption to the localized surface plasmon resonance of the Ni.
nanoparticles, which is known to have a broad band in the IR-vis range, as was previously reported [52–54].

To take advantage of the extra light absorption of the catalyst used in this work, we performed a series of CO₂ methanation tests under solar illumination. We observed a significant increase in the CH₄ yield under illumination, which was concomitant with a temperature increase in the reactor.

Although the plasmonic Ni nanoparticles have a higher damping factor comparing to frequently used Au or Ag nanoparticles, the overlap of their absorption spectrum with the solar one is much better [55]. This ensures an effective absorption of the solar illumination and transforming it to heat.

We have used three different light intensities and operated at three different starting temperatures in order to study the effects related to the photoinduced catalytic activity. Also, in order to investigate the nature of the activity enhancement, we performed wavelength-dependent tests by introducing bandpass filters into the experimental setup. UV filter (λ > 400 nm) was used to cut off the UV part of the solar spectrum, while the IR filter (λ < 800 nm) removed the near-IR and IR photons. Transmission spectra of the used filters are shown on the Fig. 6(b). We have used the combination of two bandpass filters in order to study the pure effect of visible light on the CO₂ conversion. It is worth noting that introducing of the UV filter solely does not affect the catalytic performance of the Ni/CeO₂ catalyst, indicating that the absorption of light by CeO₂ is not responsible for the increase in the catalytic activity.

Fig. 7(a) sums up the changes in catalytic activity that occur in the

| Table 1 |
|---|
| N₂-physisorption and CO-chemisorption analysis data of the Ni/CeO₂ catalyst, pure CeO₂ support and SBA-15 mesoporous silica template. |
| Sample | Surface area, m²/g | Micropore volume, cm³/g | Pore size, nm | Ni dispersion, % | Active metal surface area, m²/g |
|---|---|---|---|---|---|
| SBA-15 | 866.2 | 0.99 | 6.6 | – | – |
| CeO₂ | 121.1 | 0.21 | 3.3 | – | – |
| Ni/CeO₂ | 103.5 | 0.20 | 2.9 | 1.3 | 8.6 |

Fig. 2. HRTEM micrographs of a detail of the non-reduced NiO/CeO₂ (left) and reduced Ni/CeO₂ (right) nanostructures. In the top panel we report the magnification in the squared area and the corresponding indexed power spectra. In the bottom panel, we report the frequency filtered maps evidencing NiO, Ni and CeO₂ planes with different colors.
catalyst under solar light illumination at different temperatures, light intensities and illumination ranges. As it can be seen from the graph, introducing of visible light has a notable effect on the reaction rate, while the full solar spectrum induces an even higher increase to the catalytic activity. As it was stated previously, cutting off the UV part of the solar spectrum (\(h\nu > 3.1 \text{ eV}\)) does not affect the reaction rate, thus neglecting the role of band-band electron transitions in ceria in the methane production enhancement. Eventually, the illumination of the catalyst may increase the rate of the methanation reaction as high as 2.4 times under suitable reaction conditions (Fig. 7(b)).

### 3.4. DRIFTS characterization

To study the effect of illumination on the intermediate steps of the reaction, we have performed in-situ DRIFTS experiments at different starting temperatures. The measurements were held both in dark conditions and under solar illumination. The corresponding spectra are shown on the Fig. 8 except the band at 2350 cm\(^{-1}\) corresponding to gas phase CO\(_2\).

The pure CeO\(_2\) samples without Ni nanoclusters show presence of hydroxyl groups with vibrations \(v(\text{OH})\) identified at 3725-3560 cm\(^{-1}\) and carbonate species only (Fig. S3). At 100 °C the surface is covered by bidentate carbonates (1586, 1300 cm\(^{-1}\)), bridged carbonates (1646, 1120 cm\(^{-1}\)) and hydrogen carbonates (1681, 1632, 1392 and 1212 cm\(^{-1}\)), presumably formed over OH groups on the CeO\(_2\) surface. The temperature increase stimulates transformation of these carbonates into more stable mono- and polydentate carbonates (1483–1500, 1357 and 1150 cm\(^{-1}\)). In the absence of Ni\(^+\) the dissociation of the H\(_2\) molecule is not favored, and, therefore, formates are not formed on pure CeO\(_2\) surface.

For Ni/CeO\(_2\) catalyst at 100 °C no methane is produced, and CO\(_2\) is mainly adsorbed on the basic Ce\(^{3+}\) sites \([56,57]\), forming carbonate and hydrogen carbonates species, which are known as precursor for creation of formates. The bands at 1680 cm\(^{-1}\), 1417 cm\(^{-1}\) and 1209 cm\(^{-1}\) were attributed to hydrogen carbonates vibrations, and the peak at 1642 cm\(^{-1}\) was assigned to bridged carbonates. The broad band (1472–1530 cm\(^{-1}\)) and a peak at 1150 cm\(^{-1}\) were attributed to mono- and polydentate carbonates. At 150 °C the weakly bound bridged carbonates are being transformed into mono- and polydentate carbonates and a new bands (1583, 1371 and 1343 cm\(^{-1}\)) attributed to formates begin to grow \([58]\). Starting from 200 °C a band attributed to production of methane appears at 3016 cm\(^{-1}\). It grows further with the temperature increase, while the intensity of the formate peaks is being diminished. On the other hand, the band, attributed to mono- and polydentate carbonates, continuously grows with the temperature increase. This testifies that formates are the precursor in the methane formation, while the mono- and polydentate carbonates species do not take significant part in the reaction.

Under illumination, the DRIFTS spectra undergo some qualitative changes. At 100 °C the peaks at 1583 and 1371 cm\(^{-1}\), attributed to vibration of monodentate formate \((v_{\text{asym}}(\text{COO}))\) become much more pronounced, while the peaks related to hydrogen carbonate species (1680, 1417 or 1209 cm\(^{-1}\)) are distinctly decreased.

Another significant feature of the DRIFTS spectra, recorded under
illumination is appearance of the carbonyl species on the Ni surface, identified at 2041 cm\(^{-1}\) (monocarbonyl), 1927 cm\(^{-1}\) (bridged carbonyl) and 1830 cm\(^{-1}\) (two-fold bridged carbonyl). These bands are growing with temperature increase up to 200\(\,^{\circ}\)C and afterwards turn to decrease along with formation of methane. This evidently indicates that carbonyls take part in the process of CO\(_2\) hydrogenation to methane. The formation of carbonyls on the Ni surface may occur either through the direct dissociation of CO\(_2\) or through decomposition of formate species.

### 3.5. Mechanism of photothermal CO\(_2\) methanation

Typically, CO\(_2\) hydrogenation on the Ni-based catalysts with inactive support, such as Al\(_2\)O\(_3\) or SiO\(_2\) occurs through the dissociation mechanism, i.e. formation of carbonyl or formate intermediates on the Ni\(^0\) active sites [43]. On the other hand, Ni/CeO\(_2\) catalysts were found to favor the associative mechanism of CO\(_2\) methanation [59–61]. It means that the CO\(_2\) molecules are chemisorbed as monodentate or bidentate carbonates on the reduced active sites of CeO\(_2\), mostly present at the Ni/CeO\(_2\) interface. At the same time, H\(_2\) molecules dissociate on the metallic Ni clusters, facilitate C–O double-bond breaking and creation of the formates followed by their step-by-step hydrogenation to CH\(_4\). This mechanism of CO\(_2\) activation is energetically more favorable than the route through CO formation, and is ensured by high mobility of the oxygen vacancies in ceria. The latter allows partial reduction of the CeO\(_2\) material by H\(_2\) at the Ni/CeO\(_2\) interface with subsequent formation of water molecules, while keeping the active sites reduced.

In dark conditions, the DRIFTS data recorded from our samples are in agreement with described above associative mechanism, indicating that stable bridged formate species serve as a main precursor for CO\(_2\) hydrogenation to methane, probably through formation of formaldehyde and methoxy adsorbates. However, when the samples are illuminated, we observe immediate changes in the DRIFTS spectra. In particular, variations of the peaks suggest that hydrogen carbonates are more likely transformed to formates. Besides, we detect the appearance of carbonyl...
species, which may be formed through dissociation of excessive formates [62].

Based on the obtained results, we suggest that illumination has a two-fold effect on the operational performance of the Ni/CeO$_2$ catalyst. First, visible and IR irradiations excite LSPR in Ni nanoparticles. Although plasmons in Ni are effectively damped, they result in strong local heating of the nickel nanoclusters with following heat dissipation to the surrounding oxide matrix. Such a plasmon-induced increase of the temperature facilitates the principal reaction channel at the Ni/CeO$_2$ interface.

On the other hand, the illumination is changing the surface chemistry of the catalyst as a result of photoelectronic processes. The obtained changes in the DRIFTS spectra indicate that illumination is promoting the stabilization of formate species from less stable hydrogen carbonates.

As the illumination of the catalyst is followed by the temperature increase (summarized in the Table S1), we have further analyzed the data from Fig. 7 in order to find out the nature of the improved catalytic activity. A thermally activated methanation reaction follows an exponential trend, i.e. $k \sim e^{E_a/kT}$, where $E_a$ is the activation energy of the reaction, or the rate-limiting step [63]. Therefore, $\ln(k) \sim -\frac{E_a}{kT}$ for the

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**Fig. 8.** In-situ DRIFTS spectra measured during CO$_2$ methanation over the Ni/CeO$_2$ catalyst (a) in dark conditions; (b) under solar illumination (5 suns).

**Fig. 9.** Arrhenius slope of the temperature activated methanation reaction. The y axis indicates the natural logarithm of catalytic activity under different reaction conditions.
process that is fully controlled by the temperature. We have plotted a natural logarithm of the measured catalytic activities in all of the applied conditions versus the inverse temperature (Fig. 9). Remarkably, all the light-assisted measurements of the catalytic activity at low conversion rates were followed the same linear trend, indicating that the solar light illumination of the Ni/CeO$_2$ catalyst caused an increase of the catalytic activity due to the local temperature changes induced by plasmonic heating of the Ni nanoparticles.

From the Fig. 9 we have evaluated the Arrhenius slope for the methanation reaction (dashed line), with the activation energy of 0.96 eV. As discussed above, we attribute this energy to the methanation pathway that takes place through hydrogenation of the formates on ceria, presumably to the dissociation of formalin to formaldehyde, as it was shown in [65]. The light-induced increase in the concentration of formates, seen from the DRIFTS data, does not affect the main reaction channel, the latter being totally controlled by temperature. However, we have found that with increase of the illumination intensity the formate and carbonyl peaks on the DRIFTS spectra become more pronounced.

We applied Gaussian multi-peak fitting to the spectra measured under dark conditions, 3 and 5 suns illumination in order to quantify the contribution of the formate intermediates depending on the intensity of the photon flux (Fig. S4). We found that the amount of adsorbed species is proportional to the power of the applied illumination, which indicates that the electronic effect of the solar illumination can be much more noticeable when higher light intensities are applied.

However, when reaching higher temperatures, one can observe a strong deviation from the linear trend towards lower catalytic activities, which points out on a new rate-limiting process that occurs at higher temperatures. As the selectivity to methane was not altered upon the temperature increase (see Table S2), this deviation cannot be explained by the activation of a side reaction, i.e. producing of carbon monoxide (reverse water gas-shift reaction), which is enabled at higher temperatures [65]. It is well known that the main deactivation pathways for the Ni-based catalysts are sintering of the Ni nanoparticles and coke formation [66]. However, these processes are also taking place at higher operation temperatures. On the other hand, Cárdenas-Arenas et al. conducted isotopic experiments on Ni/CeO$_2$ catalyst and showed that water desorption is the slowest step of the methanation mechanism [61]. As it was mentioned before, the light-assisted reaction was held in a custom-designed planar-type reactor under low gas flows. Apparently, at higher CO$_2$ conversion values the physisorbed water molecules, formed in big amounts during the reaction, are not efficiently removed from the reactor and block the active sites of the catalyst. Thus, in our case, desorption of water becomes a new rate-limiting step in the methanation process at high conversion rates. We assume that changing of the reactor design to being suitable for high-flow operation conditions will allow to examine the effect of strong illumination on methanation reaction rate with no limitations due to poisoning of the active sites by water molecules.

To understand the effect of different wavelengths on the catalyst we performed a series of DRIFTS experiments under solar illumination with different optical filters. The dark spectra were subtracted from the ones recorded under illumination in order to obtain differential data (Fig. 10). The results show that the UV part of the solar spectrum has the largest effect on the growth of formate and carbonyl peaks, while the visible light has a much weaker influence on the formate concentration. In the table S3 we have summarized the relative amounts of incident light intensities, used in this work. The intensity of transmitted light with the energy sufficient to excite band-to-band transitions in CeO$_2$ (250–400 nm) sums up to 10.5 % of the total solar spectrum, which results in a non-negligible amount of photogenerated carriers in the ceria support.

It should be noted that the real temperature of the illuminated catalyst is higher than the nominal value of the starting point. To ensure that the DRIFTS changes do not occur due to temperature-induced effects, caused by the heating of the catalyst under solar light, we have compared DRIFTS spectra of the catalyst under illumination with spectra recorded at elevated temperature in dark conditions (Fig. S5). The differential graph exhibits the same features as in the Fig. 10, which proves the non-thermal nature of the light-induced changes on the catalyst surface.

According to DRIFTS data, solar light promotes stabilization of the formate species on ceria surface. The increased concentration of formates, however, does not facilitate the overall methane production at the Ni/CeO$_2$ interface, which follows the Arrhenius slope (Fig. 9).

The surface of cerium oxide is known to be enriched with oxygen vacancies, which can trap photogenerated electrons under UV light excitation [67,68]. Furthermore, it was reported that chemisorption of CO$_2$ becomes favorable on the reduced CeO$_2$ surface with abundance of additional charge [57,69]. Therefore, it is plausible that UV and blue light illumination, present in the solar spectrum, can generate electron-hole pairs in ceria that activate surface vacancy sites and enhance sorption processes at CeO$_2$ surface. Photogenerated carriers facilitate transformation of the carbonate species to the more energetically stable formates, which is reflected in the corresponding changes of DRIFTS spectra. The concentration of formates is increased throughout the ceria support, including the sites apart from the Ni/CeO$_2$ interface. On the other hand, hydrogenation of formates, being the rate-limiting step of the methanation reaction, requires active hydrogen atoms, which are abundant at the interface with nickel nanoparticles. Thus, the formates adsorbed on ceria surface apart from the interface are lacking hydrogens and do not contribute effectively to the overall methane production.

The suggested mechanism of CO$_2$ methanation on the Ni/CeO$_2$ catalyst is schematically shown on Fig. 11 and can be described as follows. In dark conditions, CO$_2$ is adsorbed on the active sites at Ni/CeO$_2$ interface as bridged carbonates or hydrogen carbonates, which are further transformed to formates and monodentate carbonates. Under constant supply of hydrogen atoms from Ni nanoclusters, the formates at the interface are hydrogenated to methane. Solar illumination has a two-fold effect on the reaction. First, Ni nanoparticles exhibit a rapid local heating due to the strongly damped localized surface plasmons generated on Ni. This leads to an increase of the average temperature of the catalyst and to the overall increase of the methane production. Secondly, UV and visible illumination is enhancing photocatalytic properties of ceria. Photoinduced charge carriers facilitate transformation of...
hydrogen carbonates to formates, giving rise to the corresponding peak on the DRIFTS spectra. However, the newly formed species adsorbed apart from the Ni/CeO$_2$ interface do not contribute to the methanation due to the limited hydrogen supply.

3.6. Energy consumption

One of the effective ways of exploiting the plasmon-induced photothermal effect of the Ni/CeO$_2$ catalyst for CO$_2$ methanation is to maintain the constant temperature of the reaction, by controlling the power supplied to the reactor. After compensation of the increased temperature to the initial value by reducing the power consumption, no improvement in the reaction rate was observed, which confirms that the photothermal effect of the illumination on the main reaction pathway is dominant at low temperatures and light intensities. However, we have measured the power, supplied to the heater of the reactor in the dark conditions ($P_{\text{dark}}$), and the corresponding power under illumination ($P_{\text{light}}$), reduced in order to maintain the working temperature at its initial value. Then, the reduced power consumption (RPC) for all of the experimental conditions used in this work was calculated according to the following formula:

$$RPC = \frac{P_{\text{dark}} - P_{\text{light}}}{P_{\text{dark}}} \times 100\%$$

The results are presented in the Fig. 12.

We can see that using concentrated sun illumination we are able to save up to 20% of power applied for the conventional thermocatalytical approach to CO$_2$ methanation. Utilization of renewable solar energy source and an earth-abundant material with plasmonic properties for facilitation of effective thermal-driven CO$_2$ methanation allowed us to gain energy for further lowering of the cost, scalability and industrialization of the process.

4. Conclusions

In summary, we have demonstrated that the catalytic activity of the methanation reaction can be improved by the photothermal effect due to localized surface plasmonic properties of the Ni nanoparticles in the Ni/CeO$_2$ catalyst under visible and infrared light illumination. Using of the mesoporous SBA-15 silica template allowed us to synthetize large-surface-area Ni/CeO$_2$ catalyst, which ensures highly stable and efficient conversion of CO$_2$ (80%) with high selectivity to methane (95%). The obtained catalyst exhibits up to a 2.4-fold increase in the reaction rate under solar light illumination, which allowed us to decrease the power consumption by 20% with respect to the dark conditions.

In-situ DRIFTS analysis revealed the mechanism of the CO$_2$ methanation in dark and light-assisted conditions. In dark conditions the CO$_2$ methanation is following the associative route through formate hydrogenation on CeO$_2$, using Ni$^0$ active sites only for H$_2$ dissociation and spillover. When illuminating the catalyst with solar light, two main effects were observed. On the one hand, the visible and IR illumination results in a strong photothermal plasmon-induced effect, followed by local heating and enhanced methane production. On the other hand, UV and blue light induced photogenerated free carriers in CeO$_2$ can improve the CO$_2$ coordination and promote the creation of formates. However, their further hydrogenation remains fully controlled by the
photothermally activated process due to the limited hydrogen supply to the formate species. Investigation of the rate-limiting step by means of the first-principles calculations will be the subject of the future work.

At high conversion rates, the activity of the Ni/CoO2 catalyst in the current reactor design is damped by the slow water desorption process from the active NiO sites. Therefore, we have utilized the plasmon-induced photothermal effect of the Ni/CoO2 catalyst for CO2 methanation by operating at low reaction temperatures, reducing the power supplied to the reactor. Nevertheless, we believe that new reactor design may open a possibility of a thorough study and exploitation of the new light-induced methanation route at higher illumination intensities and conversion rates, thus further reducing the power consumed by the reactor.

Our results emphasize the two-fold effect of solar illumination on the plasmonic Ni-based catalysts: photothermal and electronic. Hence, we expect the combined photo- and thermental approach to provide a promising pathway for Ni-based heterogeneous catalysts for green CO2 hydrogenation.

CRediT authorship contribution statement

Viktoria Golovanova: Writing - original draft, Conceptualization, Investigation. Maria Chiara Spadaro: Investigation, Writing - review & editing. Jordi Arbiol: Investigation, Writing - review & editing. Via-cheslav Golovanov: Writing - review & editing, Conceptualization. Tapio T. Ranta-la: Writing - review & editing, Teresa Andrea: Writing - review & editing, Supervision, Conceptualization. Joan Ramon Morante: Writing - review & editing, Supervision, Conceptualization.

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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