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CO₂ Hydrogenation at Atmospheric Pressure and Low Temperature Using Plasma-Enhanced Catalysis over Supported Cobalt Oxide Catalysts

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ABSTRACT: CO₂ is a promising renewable, cheap, and abundant C1 feedstock for producing valuable chemicals, such as CO and methanol. In conventional reactors, because of thermodynamic constraints, converting CO₂ to methanol requires high temperature and pressure, typically 250 °C and 20 bar. Nonthermal plasma is a better option, as it can convert CO₂ at near-ambient temperature and pressure. Adding a catalyst to such plasma setups can enhance conversion and selectivity. However, we know little about the effects of catalysts in such systems. Here, we study CO₂ hydrogenation in a dielectric barrier discharge plasma-catalysis setup under ambient conditions using MgO, γ-Al₂O₃, and a series of Co₃O₄/MgO catalysts. While all three catalyst types enhanced CO₂ conversion, Co₃O₄/MgO gave the best results, converting up to 35% of CO₂ and reaching the highest methanol yield (10%). Control experiments showed that the basic MgO support is more active than the acidic γ-Al₂O₃, and that MgO-supported cobalt oxide catalysts improve the selectivity toward methanol. The methanol yield can be tuned by changing the metal loading. Overall, our study shows the utility of plasma catalysis for CO₂ conversion under mild conditions, with the potential to reduce the energy footprint of CO₂-recycling processes.

KEYWORDS: green chemistry, plasma catalysis, methanol synthesis, cobalt oxide, CO₂ conversion

INTRODUCTION

The increasing amount of CO₂ in the atmosphere is a global problem.⁻¹⁻³ Yet CO₂ is also an important chemical resource that should not be thrown away. Ideally, we should use it as a raw material to make valuable products. This can be done via carbon capture, either from the atmosphere or from industrial flue-gases, followed by a chemical reaction.⁻⁴⁻⁶ Typically, this would involve a catalytic process. Hydrogenation is a versatile option for CO₂ valorization, provided that renewable hydrogen is used.⁻⁷⁻¹¹ One can alter the reaction parameters (CO₂/H₂ ratio, temperature, and pressure) and the catalyst composition to control the product distribution.¹² CO₂ can be hydrogenated to carbon monoxide (CO), methane (CH₄), methanol (CH₃OH), ethanol (C₂H₅OH), and lower olefins (C₃⁻C₅). Of these, methanol is especially interesting, as it is a precursor for formaldehyde, dimethyl ether, gasoline, and olefins.¹³⁻¹⁴ Methanol is also an efficient energy carrier.¹⁵⁻¹⁷ CO₂ hydrogenation to methanol (eq 1) is exothermic and, therefore, favored at low temperatures. This reaction is also favored at high pressures because fewer molecules are produced. However, the high thermodynamic stability of CO₂ (ΔG° = −394.4 kJ mol⁻¹) requires high temperatures for high conversion. The reverse-water gas shift reaction (RWGS, eq 2) is favored at < 200 °C because of its endothermic character. The side product, CO, is itself a useful building block for making a variety of chemicals.¹²⁻¹⁸ Therefore, CO₂ hydrogenation to methanol is typically performed at 250 °C and 20 bar.¹⁴⁻¹⁹

\[
\text{CO}_2 + 3\text{H}_2 \leftrightarrow \text{CH}_3\text{OH} + \text{H}_2\text{O} \quad \Delta H_{\text{298 K}} = -49.5 \text{ kJ mol}^{-1} \tag{1}
\]

\[
\text{CO}_2 + \text{H}_2 \leftrightarrow \text{CO} + \text{H}_2\text{O} \quad \Delta H_{\text{298 K}} = 41.2 \text{ kJ mol}^{-1} \tag{2}
\]

Yet there is an alternative to thermal catalysis: plasma-enhanced catalysis can be used for converting CO₂ to useful chemicals at near-ambient temperatures and pressures.⁻²⁰⁻²⁹ In nonthermal plasmas, high-energy electrons (with an average electron temperature of 1–10 eV) collide with stable molecules and activate them, while the bulk gas temperature remains low.³⁰⁻³¹ Catalysts can be introduced into these systems to increase the conversion and control the
selectivity.\textsuperscript{19,32} The most common option in plasma catalysis is the dielectric barrier discharge (DBD), among the different types of plasma discharges. This is because DBD reactors usually run at low temperatures and atmospheric pressure, reducing operating costs and complexity, and allow facile combination with catalysts.\textsuperscript{30,33,34} Despite the growing research activity in plasma-enhanced catalysis, we still do not fully understand the relationship between the catalyst properties and overall reaction performance.\textsuperscript{35–38} This is because of the variety of the reactions and plasma configurations, which lead to different plasma species and different plasma–catalyst interactions.\textsuperscript{19–42} Here, we study the effect of basic catalysts (MgO and CoO\textsubscript{x}/MgO catalysts) on conversion and product selectivity during CO\textsubscript{2} hydrogenation. We ran the reaction in a water-cooled DBD plasma-catalysis setup, at 35 °C and ambient pressure. Furthermore, we tested γ-Al\textsubscript{2}O\textsubscript{3} to check if the enhanced adsorption of acidic CO\textsubscript{2} on basic supports is also valid under plasma conditions.\textsuperscript{43} We observed that basic materials enhance CO\textsubscript{2} conversion, and that the production of methanol is related to the metal loading and the dispersion of metal–support interface sites in CoO\textsubscript{x}/MgO catalysts.

## EXPERIMENTAL SECTION

### Materials and Instrumentation.
Powder X-ray diffraction (XRD) analyses were carried out on a MiniFlex II diffractometer using Cu Kα radiation (X-ray tube set at 30 kV and 15 mA). The XRD patterns were recorded between 2θ = 20–80° at a speed of 2.5°/min.\textsuperscript{4} A Thermo Scientific Surfer instrument was used to carry out N\textsubscript{2} adsorption–desorption analyses at 77 K. The samples were pretreated under vacuum at 200 °C for 6 h. Surface areas were determined with the Brunauer–Emmett–Teller (BET) method, and the mesoporosity was analyzed using the Barrett, Joyner, and Halenda (BJH) method. Hydrogen temperature-programmed reduction (H\textsubscript{2}-TPR) profiles were obtained using a TPDRO Series 1100 from Thermo Scientific, following the procedure previously reported by Ronda-Lloret et al.\textsuperscript{6} High-resolution transmission electron microscopy (HRTEM) micrographs and transmission electron microscopy coupled with energy-dispersive X-ray spectroscopy (STEM–EDS) images were obtained on a JEOL-JEM 2100F microscope running at 200 kV. X-ray photoelectron spectroscopy (XPS) analyses were performed with a SPECS PHIIBOS 100 MCDS hemispherical electron analyzer operating in a constant pass energy. The analysis details were described previously in Matthaiou et al.\textsuperscript{44}

### Procedure for Catalyst Synthesis.
Using Co(NO\textsubscript{3})\textsubscript{2}·6H\textsubscript{2}O (99%, Acros Organics) as the metal oxide precursor and MgO (Sigma-Aldrich) as the support, we prepared samples containing different percentages of cobalt (5, 10, 15, and 20 wt % on Co metal-basis) by wet impregnation.\textsuperscript{4} After impregnation, the materials were dried at 120 °C for 2 h and then calcined in a muffle furnace at 450 °C for 4 h. Bulk CoO\textsubscript{x} was prepared by mixing Na\textsubscript{2}CO\textsubscript{3}, polyethylene glycol, and Co(NO\textsubscript{3})\textsubscript{2}·6H\textsubscript{2}O in water, as reported earlier.\textsuperscript{45} After stirring for 3 h at room temperature, the precipitate was centrifuged and washed several times with water and ethanol. After drying at 65 °C for 6 h in a vacuum oven, the sample was calcined in air at 300 °C for 3 h.

#### Plasma Setup and Plasma-Catalytic Tests.
The plasma-catalytic tests were performed in a coaxial DBD reactor with a 50 mm discharge length and a 1 mm discharge gap (Scheme 1). Circulating water was used as the ground electrode and cooling system (Grant LT Ecocool 150) to keep the reaction temperature at 35 °C. An AC high-voltage power supply (with a peak-to-peak voltage of up to 30 kV) was used to ignite the plasma with a fixed frequency of 9.2 kHz. CO\textsubscript{2} and H\textsubscript{2} (H\textsubscript{2}/CO\textsubscript{2} = 3:1) were used as reactants with a total flow rate of 28 mL·min\textsuperscript{-1}. The catalysts (500 mg) were fully packed in the discharge area, the plasma reaction was run for 1.5 h and the products were then analyzed.
The characteristic peaks of Co₃O₄ and CoO are not visible, as they overlap with those of MgO. HRTEM and STEM images of 15% Co₃O₄/MgO catalyst do not follow this trend, indicating the presence of larger cobalt oxide particles that block the pores and decrease the surface area and pore volume values increase with the metal loading up to 15% (Table 1). 20% Co₃O₄/MgO catalyst does not follow this trend, indicating the presence of larger cobalt oxide particles that block the pores and decrease the surface area and pore volume values increase with the metal loading up to 15% (Table 1).

**RESULTS AND DISCUSSION**

**Catalyst Synthesis.** We tested MgO and γ-Al₂O₃ as packing materials in the DBD plasma. MgO was also used as support for the cobalt oxide catalysts with different cobalt metal loadings. We also prepared bulk Co₃O₄ by mixing polyethylene glycol and cobalt nitrate hexahydrate in water (see the detailed procedures in the Experimental Section).

**Catalyst Characterization.** The XRD patterns of the fresh Co₃O₄/MgO catalysts show that cobalt oxide nanoparticles are highly dispersed over the support (Figures 1 and 2). The average size of cobalt oxide nanoparticles was 27.7 ± 11.5 nm. Moreover, HRTEM shows lattice fringes with interplanar distances of 0.24 and 0.28−0.29 nm (Figure 1b,c), assigned to the (311) and (220) planes of cobalt oxide nanoparticles, respectively. The selected-area electron diffraction (SAED) pattern shows diffraction rings characteristic of a polycrystalline material, corresponding to the MgO [(220), (200), and (311)] and cobalt oxide nanoparticles (311) planes.

We then used XPS to study the surface composition of the catalysts. The Co 2p spectra of the Co₃O₄/MgO catalysts show the doublet of two spin−orbit components, Co 2p₃/2 and Co 2p₁/2 (Figure 3). The relative atomic percentage of Co₃O₄ species is higher than that of CoO species for all fresh catalysts, indicating that the surface of the catalysts mainly contains Co₃O₄ (Table S1). This reveals that Co₃O₄−CoO/MgO interface sites are available on the catalyst surface as active centers for CO₂ hydrogenation. The O 1s, C 1s, and Mg 2p spectra of the 15% Co₃O₄/MgO catalyst, as well as details of the binding energies are included in the Supporting Information (Figure S2 and Tables S2−S5).

The H₂-TPR profiles of the as-prepared materials (Figure S3, bottom profiles) show two regions. The low-temperature region (150−450 °C) is assigned to the reduction of “free” Co₃O₄ (supported cobalt oxide, i.e., interacting weakly with the support, thus not forming a stable compound). This reduction peak contains two or more contributions, corresponding to the Co₃O₄ step wise reduction sequence where Co₃O₄ is reduced to CoO, and CoO is reduced to metallic cobalt at higher temperatures. The reduction of Co₃O₄ species with different interaction with the support can also lead to different contributions in the reduction peak. They are less defined in the 20% Co₃O₄/MgO sample, most likely because the first contribution shifts to a higher temperature. This reflects the presence of larger cobalt oxide particles, which are harder to reduce than smaller ones.

In the high temperature region (500−1000 °C), the 5% Co₃O₄/MgO reduction profile clearly shows a small peak at 500−600 °C, ascribed to the reduction of MgCo₂O₄ species. Above 700 °C, all the samples show the reduction of stable (Co,Mg)O solid solution species. These peaks overlap at higher metal loadings. In some cases, the (Co,Mg)O solid solution is only partially reduced.

The fresh catalysts showed nitrogen adsorption−desorption isotherms (Figure S4) similar to type IV, with a hysteresis feature characteristic of mesoporous materials. The BET surface area and pore volume values increase with the metal loading up to 15% (Table 1). 20% Co₃O₄/MgO catalyst does not follow this trend, indicating the presence of larger cobalt oxide particles that block the pores and decrease the surface energy efficiency (mmol kWh⁻¹)

\[ \text{energy efficiency} = \frac{\text{converted product (mmol h⁻¹)}}{\text{discharge power (W)}} \]
HRTEM analysis of this sample confirmed the presence of larger cobalt oxide particles compared to the 15% sample (Figure S5a).

**Catalytic Tests.** We performed the CO\(_2\) hydrogenation reaction in a coaxial DBD reactor, maintained at 35 °C using circulating cooling water. The plasma operated at atmospheric pressure and 10 W. Previous plasma-catalytic CO\(_2\) hydrogenation studies show that an excess of hydrogen in the reaction mixture, up to a 4:1 H\(_2\)/CO\(_2\) molar ratio, typically increases the conversion.\(^{22,25,58}\) In addition, two recent studies reported higher methanol yields at a 3:1 H\(_2\)/CO\(_2\) molar ratio.\(^{26,27}\) To favor CO\(_2\) conversion and methanol formation, we set the H\(_2\)/CO\(_2\) molar ratio to 3:1.

First, we tested plasma alone (Figure S6), which gave 15% CO\(_2\) conversion and 7% H\(_2\) conversion (the plasma background reaction). The products were mainly CO (71% selectivity) and methanol (22%), as well as 2% methane and 6% acetic acid (all results are averages of triplicate measurements).

We then tested the effect of different packing materials. MgO and \(\gamma\)-Al\(_2\)O\(_3\) are conventional supports, which are usually inactive in thermal catalysis. However, these materials enhanced the activity when introduced in the DBD plasma chamber even under the mild reaction conditions mentioned above (Figure S6). MgO showed significantly higher conversion values (27% CO\(_2\) conversion and 13% H\(_2\) conversion) than plasma alone. CO selectivity also increased, from 71% with plasma only to 84% with MgO. Alumina also gave higher CO\(_2\) and H\(_2\) conversion (17 and 7%, respectively) than plasma alone, and higher CO selectivity (80%).

The changes in conversion and product distribution when placing a packing material in the plasma discharge can be explained by examining the possible reaction pathways. Typically, in a CO\(_2\)/H\(_2\) mixture in DBD plasma (without catalyst), consumption of CO\(_2\) predominantly occurs via electron impact dissociation to CO (R1).\(^{59,60}\) The contribution from the electron impact vibrational excitation of CO\(_2\) is negligible because of its tendency to react back to ground CO\(_2\). Similarly, H\(_2\) is consumed by electron impact dissociation to two hydrogen radicals (R2). CO and H\(^+\) radicals can start several reactions that lead to formaldehyde CH\(_2\)O (R3–R9). CH\(_2\)O is then hydrogenated to methanol (R11 and R12),\(^{26,59}\) while water is also produced (R12 and R13).\(^{59}\)

![Figure 3. Co 2p XPS spectra of the fresh Co\(_x\)O\(_y\)/MgO catalysts.](image)

### Table 1. BET Surface Area, Cumulative Pore Volume, and Average Pore Diameter (from BJH Analysis) Values Derived from Nitrogen Sorption Isotherms

<table>
<thead>
<tr>
<th>catalyst</th>
<th>(S_{\text{BET}}) (m(^2)·g(^{-1}))</th>
<th>pore volume (cm(^3)·g(^{-1}))</th>
<th>average pore diameter (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>5% Co(_x)O(_y)/MgO</td>
<td>33</td>
<td>0.2</td>
<td>56</td>
</tr>
<tr>
<td>10% Co(_x)O(_y)/MgO</td>
<td>37</td>
<td>0.4</td>
<td>44</td>
</tr>
<tr>
<td>15% Co(_x)O(_y)/MgO</td>
<td>39</td>
<td>0.6</td>
<td>49</td>
</tr>
<tr>
<td>20% Co(_x)O(_y)/MgO</td>
<td>26</td>
<td>0.3</td>
<td>26</td>
</tr>
</tbody>
</table>
When combining plasma and a catalyst, previous work found that both plasma-assisted surface reactions and gas-phase reactions contribute to the conversion and selectivity.61 Gas-phase reactions involve reactions R1 to R11. In addition, CO derived from CO2 dissociation in the gas phase can adsorb on the catalyst surface and react with adsorbed hydrogen to form methanol, as represented in Figure 4a.61–63 CO2 can also adsorb on the catalyst surface, as confirmed elsewhere with the detection of carbonates using in situ spectroscopy analysis.23,61,62 Both ground and excited CO2 can be adsorbed, although the contribution of excited CO2 can be neglected because it is prone to quenching on the surface.26,64

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\[
\begin{align*}
\text{CO}_2 + e^- & \rightarrow \text{CO} + O^+ + e^- \\
H_2 + e^- & \rightarrow H^+ + H^+ + e^- \\
\text{CO} + H^+ & \rightarrow \text{CHO}^+ \\
\text{CHO}^+ + \text{CHO}^+ & \rightarrow \text{CH}_3\text{O} + \text{CO} \\
\text{CO} + e^- & \rightarrow C^+ + O^+ + e^- \\
C^+ + H^+ & \rightarrow \text{CH}^+ + e^- \\
\text{CH}^+ + H^+ & \rightarrow \text{CH}_2^+ \\
\text{CH}_2^+ + O^+ & \rightarrow \text{CH}_3\text{O} \\
\text{CO}_2 + \text{CH}_2 & \rightarrow \text{CH}_2\text{O} + \text{CO} \\
\text{CH}_2\text{O} + H^+ & \rightarrow \text{CH}_2\text{OH}^+ \\
\text{CH}_2\text{OH}^+ + H^+ & \rightarrow \text{CH}_3\text{OH} \\
O^+ + H^+ & \rightarrow \text{OH}^+ + e^- \\
\text{OH}^+ + H_2 & \rightarrow \text{H}_2\text{O} + H^+ \\
\end{align*}
\]
Adsorbed CO₂ is then hydrogenated to either hydrocarboxyl (HOCO) or formate (HCOO⁻) species (Figure 4b). The decomposition of HOCO to CO initiates the so-called RWGS + CO-hydro pathway. The formate pathway can also take place, where HCOO⁻ is hydrogenated in several steps. Both pathways lead to the formation of H₂CO, which is hydrogenated to methanol via the methoxy (H₃CO) intermediate.

The increase in CO₂ conversion and CO selectivity when placing MgO or Al₂O₃ in the plasma chamber is ascribed to their ability to adsorb CO₂ and perform both plasma-assisted surface reactions and gas-phase reactions. MgO shows higher conversion than Al₂O₃, which is related to their acid–base properties. CO₂ is attracted better by the basic sites of solid surfaces because of its acidic property. MgO is significantly more basic than Al₂O₃ and therefore it has a higher CO₂ adsorption capacity. Both materials increase CO selectivity, indicating that they promote the RWGS pathway. These metal oxides do not have a strong affinity to CO and H₂/H₂O species, hindering the formation of methanol.

Shifting the selectivity toward methanol requires a stronger binding energy of CO and hydrogen, as well as the participation of a considerable amount of adsorbed hydrogen in the reaction pathway. Cobalt-based catalysts are active in CO₂ hydrogenation to ethanol, methane, and larger hydrocarbons. Therefore, we tested the influence of MgO-supported cobalt oxide catalysts with different metal loadings on the reaction performance (Figure 5). Our results show an improved conversion when using CoₓOₓ/MgO catalysts, which is partially ascribed to their ability to promote surface reactions. These catalysts also gave a narrower product distribution. We also observed acetic acid (<1%), ethanol (trace), and methane (<2%).

The 5% CoₓOₓ/MgO catalyst converted 30% of CO₂ and 13% of H₂, giving 92% selectivity to CO and 6% of methanol. Compared to MgO, it only improved CO₂ conversion, thereby increasing CO selectivity. The low amount of cobalt oxide in this catalyst does not shift the product distribution to methanol.

The 10% CoₓOₓ/MgO catalyst showed 31% CO₂ conversion and 15% H₂ conversion, as well as 76% selectivity to CO and 20% selectivity to methanol. Further increasing the cobalt loading to 15% improved the conversion values to 33% CO₂ and 24% H₂, and the methanol selectivity to 31%. This catalyst gave the highest methanol yield (>10%). This indicates that catalysts with high metal loading increase the amount of adsorbed CO and hydrogen available for the reaction, facilitating methanol production. Increasing the metal loading to 20% gave 31% CO₂ conversion and 9% methanol yield.

Figure 5. CO₂ and H₂ conversion, selectivity and methanol yield plots of plasma alone, MgO, CoₓOₓ/MgO with different cobalt metal loadings, and unsupported Co₃O₄. Reaction conditions: 35 °C, atmospheric pressure, 500 mg catalyst, a total flow of 28 mL·min⁻¹, a H₂/CO₂ molar ratio of 3:1, a discharge power of 10 W, and a reaction time of 1.5 h.
From these results, we conclude that the quantity and dispersion of the Co$_3$O$_4$/CoO sites play an important role in the activity enhancement and methanol selectivity, in agreement with previous studies.\textsuperscript{65,73} CO$_2$ conversion and methanol selectivity increase when increasing the metal loading from 5 to 15\%. The BET surface area and pore volume also increase when increasing the metal loading up to 15\% (see Table 1). This indicates that a good dispersion of the metal oxide nanoparticles on the support favors CO$_2$ conversion. Higher metal loadings (20\%) lead to larger cobalt oxide particles (as confirmed by HRTEM), which block the pores of the support and decrease the catalyst surface area. This reduces the metal oxide dispersion compared to lower metal loadings, inhibiting a further increase on conversion and methanol production.

For comparison, we tested unsupported Co$_3$O$_4$ (Figure 5). This gave lower conversion (8\% CO$_2$ and 10\% H$_2$) than the supported Co$_{x}$O$_{y}$/MgO catalysts, with CO as the main product (98\% selectivity). The decrease in methanol formation when using Co$_3$O$_4$ emphasizes the importance of metal oxide-support interface sites for methanol synthesis.

Compared to plasma only, the combination of plasma and any tested catalyst improves the energy efficiency of CO production (Figure S7), reaching its maximum when using 10\% Co$_{x}$O$_{y}$/MgO catalyst (509 mmol CO-kWh$^{-1}$). The energy efficiency of methanol production is improved when a considerable metal loading of cobalt is used (10, 15, and 20\% Co$_{x}$O$_{y}$/MgO catalysts). The optimal methanol energy efficiency was obtained with 15\% Co$_{x}$O$_{y}$/MgO (190.8 mmol MeOH-kWh$^{-1}$), which is related to the balance between a considerable amount of Co$_3$O$_4$/CoO sites and their good dispersion on the support. None of these catalysts were active in thermal activity tests, showing the importance of the plasma-catalytic hybrid approach in activating CO$_2$ at near-ambient temperatures. In addition, we studied the stability of the 15\% Co$_{x}$O$_{y}$/MgO catalyst under reaction conditions (Figure S8). The catalyst performance was stable up to 4.5 h, with only a slight deactivation at the start of the reaction.

**Plasma and Catalyst Interactions.** We studied the effect of the plasma discharge on the physicochemical properties of the Co$_{x}$O$_{y}$/MgO catalysts. The XRD patterns of the spent catalysts correspond to those of the fresh catalysts (Figure S1b). HRTEM and STEM images of 15\% Co$_{x}$O$_{y}$/MgO spent catalyst (Figures S8 and S9) discard the sintering of cobalt oxide particles because the average size of cobalt oxide nanoparticles (24.5 ± 7.6 nm) is very similar to the fresh catalyst (27.7 ± 11.5 nm). HRTEM images of spent 20\% Co$_{x}$O$_{y}$/MgO (Figure S5b) do not show a significant change on the particle size neither. In general, the surface area and porosity of the spent catalysts are similar to the fresh ones (Table S7). The 20\% sample shows the largest increase in the BET surface area (from 26 to 35 m$^2$·g$^{-1}$), and the 15\% sample shows the largest increase in the pore volume and average pore size. In addition, XPS analysis of all catalysts shows that the $I_{Co}/I_{Mg}$ ratio remains unaltered (Table S6), indicating that the

![Figure 6. Co 2p XPS spectra of the spent Co$_{x}$O$_{y}$/MgO catalysts.](https://dx.doi.org/10.1021/acssuschemeng.0c05565)
dispersion of cobalt oxide on the support is not affected by the plasma.

XPS and H2-TPR analyses of the spent catalysts indicate that CoO/MgO surface species undergoes partial reduction under CO2/H2 plasma. For the samples with a cobalt metal loading lower than 20%, the XPS Co 2p peaks corresponding to CoO species increase in intensity, while the ones of Co3O4 decrease (Figure 6 and Table S1). The 20% CoO/MgO catalyst does not show a major reduction after plasma exposure, as the Co3O4 and CoO peak intensities remain similar to the fresh catalyst. In the reduction profiles (Figure S3), the first contribution in the low temperature region becomes less intense for the spent 5, 10, and 15% Co3O4/MgO catalysts. This contribution corresponds to the easily reducible CoO species. There are no major changes in the reduction profile of 20% Co3O4/MgO, indicating that this sample contains larger CoO particles that are harder to reduce.

We conclude that the catalysts with a cobalt metal loading lower than 20% are significantly reduced from Co3O4 to CoO in the hydrogen-rich plasma environment. Nevertheless, we cannot exclude the formation of metallic cobalt because of plasma exposure. Because all spent catalysts were stored in air prior to characterization, we cannot detect metallic cobalt in XPS and H2-TPR analysis.

We also studied the effect of the catalysts on the plasma discharge properties, which in turn might affect the reaction performance. Figure S10a shows the current signals of the plasma in the presence and absence of packing materials. The intensity and density of the current pulses of the discharge follow the order of 15% CoO/MgO > plasma alone > MgO > Co3O4, which agrees with the order of CO2 conversion. Packing MgO or Co3O4 into the discharge gap weakens the formation of microdischarges compared to plasma alone. The 15% Co3O4/MgO catalyst enhances the discharge, as evidenced by the increase in the current pulse, thereby favoring the formation of radicals and vibrationally excited species that can participate in the reaction.40,52,76 In addition, the Q−V Lissajous plot (Figure S10b) of the plasma with a packing material is very similar to the one of plasma alone, remaining with an elliptical shape. We find that the change in the cobalt loading of the Co3O4/MgO catalysts does not affect the discharge, as the electrical signals are almost the same. Thus, we attribute the difference in product selectivity when using packing materials to their properties and to the plasma-enhanced promotion of surface reactions on the metal–support interface sites (see Figure S above).

**CONCLUSIONS**

In this study, we show the potential of plasma-enhanced catalysis for CO2 hydrogenation reactions under ambient conditions. Packing a catalyst in a DBD plasma enhances the conversion and narrows the product distribution of CO2 hydrogenation. We found that the basicity of the MgO support enhances the conversion of CO2 compared to more acidic supports (γ-Al2O3). Both type of supports promote the production of CO as the main product. When testing the CoO/MgO catalysts, the quantity and dispersion of metal–support interface sites are important for the activity enhancement and methanol selectivity. Although CO is the main product, the catalysts with higher cobalt metal loadings and a good cobalt oxide dispersion favor methanol production. The most active catalyst was 15 wt % CoO/MgO, which converted 33% CO2 and 24% H2 near room temperature and at atmospheric pressure. This catalyst gave 10% methanol yield, the highest among the tested materials. A further increase in the metal loading leads to larger cobalt oxide particles that block the pores, giving lower CO conversion and less methanol. Finding correlations between catalyst properties and reaction performance remains a challenge in this field. We hope that this study will help in the rational design of materials for plasma-enhanced catalysis applications.

**ASSOCIATED CONTENT**

**Supporting Information**

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acssuschemeng.0c05565.

XRD patterns of the fresh and spent catalysts; XPS data of the fresh and spent catalysts; temperature-programmed reduction profiles of the fresh and spent catalysts; nitrogen adsorption−desorption isotherms of the fresh catalysts; HRTEM and STEM−EDS images of the spent 15% CoO/MgO catalyst; HRTEM images of the fresh and spent 20% CoO2/MgO catalyst; plasma-catalytic tests of plasma alone, MgO, and γ-Al2O3; energy efficiency values of all catalysts; stability test of the 15% CoO2/MgO catalyst; plasma electrical diagnostics (current signals, Q−V Lissajous plots, and calculated discharge properties) of plasma alone, MgO, 15% CoO2/MgO, and Co3O4 (PDF)

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**Author Contributions**

M.R.-L. and Y.W. contributed equally for this article. This article was written through contributions of all authors. All authors have given approval to the final version of the manuscript.

**Notes**

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