Nanostructure of nickel-promoted indium oxide catalysts drives selectivity in CO₂ hydrogenation

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Metal promotion in heterogeneous catalysis requires nanoscale-precision architectures to attain maximized and durable benefits. Herein, we unravel the complex interplay between nanostructure and product selectivity of nickel-promoted In₂O₃ in CO₂ hydrogenation to methanol through in-depth characterization, theoretical simulations, and kinetic analyses. Up to 10 wt.% nickel, InNi₃ patches are formed on the oxide surface, which cannot activate CO₂ but boost methanol production supplying neutral hydrogen species. Since protons and hydrides generated on In₂O₃ drive methanol synthesis rather than the reverse water-gas shift but radicals foster both reactions, nickel-lean catalysts featuring nanometric alloy layers provide a favorable balance between charged and neutral hydrogen species. For nickel contents >10 wt.%, extended InNi₃ structures favor CO production and metallic nickel additionally present produces some methane. This study marks a step ahead towards green methanol synthesis and uncovers chemistry aspects of nickel that shall spark inspiration for other catalytic applications.
In heterogeneous catalysis, numerous systems rely on metal promotion to maximize process throughput\textsuperscript{1,2}. Since these additives might carry stand-alone activity for undesired reactions jeopardizing selectivity and/or contribute to catalyst deactivation in conventional forms such as supported nanoparticles, specific nanostructures need to be designed to stabilize metal speciations displaying tailored electronic and geometric properties that minimize drawbacks while preserving benefits\textsuperscript{3-7}. This often is a challenging task since the uniform production and in-depth characterization of precise atom-resolved structures lie at the frontier of current technologies.

In the frame of mitigating global environmental changes and lessening our reliance on fossil feedstocks\textsuperscript{8-11}, indium oxide was introduced as a breakthrough catalyst for CO\textsubscript{2} hydrogenation to methanol\textsuperscript{12}, exhibiting extraordinary high selectivity and superior activity and stability when supported on monoclinic ZrO\textsubscript{2}\textsuperscript{13-15}. Mechanistic investigations indicated that vacancies formed at a specific surface lattice position mediate CO\textsubscript{2} activation and H\textsubscript{2} heterolytic splitting\textsuperscript{16-19}, the latter unlocking the preferential formation of methanol instead of CO via the reverse water-gas shift (RWGS) reaction\textsuperscript{18}. Still, since heterolytic H\textsubscript{2} activation is energetically demanding and limits the methanol synthesis rate, promotion with various hydrogenation elements was explored\textsuperscript{20-30}.

Platinum and palladium nanoparticles were shown to boost catalyst performance by aiding H\textsubscript{2} splitting, thus fostering CO\textsubscript{2} hydrogenation and generating additional vacancies on In\textsubscript{2}O\textsubscript{3}, but led to inferior methanol selectivity due to intrinsic RWGS activity and substantial reduction-induced In\textsubscript{2}O\textsubscript{3} sintering. These pitfalls were mitigated by anchoring low-nuclearity (ca. 3 atoms) clusters to the In\textsubscript{2}O\textsubscript{3} lattice in the case of gold and palladium\textsuperscript{29,31}. Ruthenium and cobalt, typical metals leading to methane, were successfully employed when the fist was alloyed with indium and the second encapsulated with In\textsubscript{2}O\textsubscript{3}\textsuperscript{23,28}. In view of these findings, the low-cost alternative nickel, also a prototypic methanation metal\textsuperscript{32}, could find effective application in CO\textsubscript{2} hydrogenation if a favorable structure is identified. A recent study showed that metallic nickel on In\textsubscript{2}O\textsubscript{3} did not produce methane in CO\textsubscript{2}-based methanol synthesis, but clear support to its segregation from the oxide was not provided\textsuperscript{26}. Another work indicated that CO\textsubscript{2} hydrogenation on Ni catalysts can be driven to CO and methanol by doping with indium\textsuperscript{35}. In methane dry reforming on In\textsubscript{2}Ni\textsubscript{x} catalysts\textsuperscript{34,35}, CO adsorption on nickel was fully suppressed upon doping with indium, suggesting their alloying. In CO\textsubscript{2} dry reforming on InNi/SiO\textsubscript{2}, the surface of Ni–In alloy particles was progressively covered with InO\textsubscript{x}H\textsubscript{y} upon use\textsuperscript{36}, as for cobalt-promoted In\textsubscript{2}O\textsubscript{3}. These radical deviations of the behavior of pure nickel from its characteristic chemistry call for a rationalization of the nanostructures underpinning them, in comparison to the other metal promoters.

Herein, the synergistic interaction of nickel with indium oxide in CO\textsubscript{2}-based methanol synthesis was explored through a comprehensive experimental and theoretical program. The nickel speciation was studied contrasting the behavior of coprecipitated and impregnated catalysts upon testing under industrially relevant conditions, with electronic effects being further evaluated on nickel deposited onto In–Al mixed oxides with distinct indium content. The nickel content was varied on the better performing impregnated catalysts identifying a clear impact on the product selectivity. In situ spectroscopy and diffraction methods along with thermal, volumetric, and microscopy analyses uncovered the nanometric construction of the selective nickel-poor and the unselective nickel-rich samples. Density functional theory (DFT) sheds light on the unique structural rearrangements of nickel deposited on In\textsubscript{2}O\textsubscript{3} and the reactivity of complementary promoted surfaces, which was linked to experimental kinetic parameters. Alloying of nickel with In\textsubscript{2}O\textsubscript{3} emerged as key to provide uncharged hydrogen atoms to active sites on In\textsubscript{2}O\textsubscript{3} while curtailing the nickel-mediated detrimental methanation pathway. Overall, this work gathers a fundamental understanding of a relevant system for sustainable methanol production and unravels structural and electronic features at the basis of the tunable selectivity of nickel in CO\textsubscript{2} hydrogenation routes.

### Results

#### Impact of nickel content and synthesis method on activity and selectivity

Nickel was incorporated into In\textsubscript{2}O\textsubscript{3} by dry impregnation (DI, 1–20 wt.%, coded as xNi-In\textsubscript{2}O\textsubscript{3}, x = 1–20) and coprecipitation (CP, 1–2.5 wt.%) aiming at a deposition on the oxide surface and formation of solid solutions to appreciate the role of metals intermixing and of the nickel oxidation state and chemical environment. N\textsubscript{2} sorption and X-ray fluorescence spectroscopy (XRF) indicated that all catalysts featured sufficiently high surface areas and nominal metal contents were closely matched (Supplementary Table 1). Assessing the samples containing 1 wt.% of nickel in CO\textsubscript{2} hydrogenation evidenced a higher methanol space-time yield (STY) compared to pure In\textsubscript{2}O\textsubscript{3} (STY = 0.16 g\textsubscript{MeOH} h\textsuperscript{−1} g\textsubscript{cat}\textsuperscript{−1}, Fig. 1a), which, after 65-h equilibration, remained at a ca. twofold higher value (0.34 g\textsubscript{MeOH} h\textsuperscript{−1} g\textsubscript{cat}\textsuperscript{−1}) for the DI catalyst and levelled to a 25% higher value (0.20 g\textsubscript{MeOH} h\textsuperscript{−1} g\textsubscript{cat}\textsuperscript{−1}) for the CP catalyst. The stable behavior of the former contrasts palladium-promoted In\textsubscript{2}O\textsubscript{3} prepared through DI, which experienced fast deactivation (Supplementary Fig. 1)\textsuperscript{31}. The inferior performance of the CP catalyst is likely due to the synthesis approach burying a substantial portion of the promoter within the bulk of In\textsubscript{2}O\textsubscript{3}, but the presence of more nickel added by this method did not yield materials

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**Fig. 1** Catalytic performance of nickel-promoted In\textsubscript{2}O\textsubscript{3} catalysts. a Space-time yield (STY) of methanol as a function of time-on-stream (TOS) over Ni-In\textsubscript{2}O\textsubscript{3} containing 1 wt.% of nickel incorporated by dry impregnation (DI) and coprecipitation (CP) and unpromoted In\textsubscript{2}O\textsubscript{3} as a reference. b CO\textsubscript{2} conversion and product selectivity of catalysts produced by DI as a function of the nominal nickel content on In\textsubscript{2}O\textsubscript{3}. Reaction conditions: T = 553 K, P = 5 MPa, molar H\textsubscript{2}:CO\textsubscript{2} = 4, and WHSV = 24,000 cm\textsuperscript{3} STP h\textsuperscript{−1} g\textsubscript{cat}\textsuperscript{−1}. 

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superior to those produced by DI (Supplementary Fig. 2). Considering the more favorable DI synthesis (Fig. 1b), methanol formation was progressively lowered in favor of the RWGS reaction up to a nickel content of 20 wt.%, while the CO₂ conversion was practically unchanged at ca. 8%, suggesting that sites activating CO₂ are unlikely located on the nickel phase. Methane formation was observed only at higher promoter contents (S₃CI₄ = 6 and 13% at 15 and 20 wt.% nickel, respectively). This hints that In₂O₃ exerts a strong influence on the nickel properties and, only for high contents, a fraction of promoter remains unperturbed and can express its intrinsic behavior. To further address the selectivity switch, the amount of indium required to trigger it was explored by supporting 5 wt.% nickel on mixed indium-aluminum oxides with variable indium content (0–100 mol% In, Supplementary Fig. 3). While nickel on pure alumina was highly selective to methane (S₃Cl₄ > 98%), 25 mol% of indium in the support sufficed to suppress methanation almost entirely (S₃Cl₄ = 3%). At the same time, the CO₂ conversion significantly dropped (from XCO₂ = 18 to 4%), corroborating that indium-modulated nickel species are significantly less active compared to pure nickel.

Characterization of nickel ‘speciation’. In-depth investigations were conducted to rationalize the behavior of the DI systems. Concerning catalyst reducibility, temperature-programmed reduction with hydrogen (H₂-TPR, Fig. 2a, Supplementary Table 2) evidenced that NiO conversion into metallic nickel occurs at ca. 340 K in all samples, preceding surface In₂O₃ reduction (370 K), which is substantially facilitated compared to the pure oxide (521 K). Based on the signal intensity in the profile of the 5Ni-Al₂O₃ reference, nickel was fully reduced in all cases. In 1Ni-In₂O₃, some In₂O₃ still reduced at its standard temperature, likely because the nickel amount was insufficient to facilitate reduction of the entire In₂O₃ surface. Diffuse reflectance infrared Fourier transform spectroscopy of adsorbed carbon monoxide (CO-DRIFTS) evidenced a weak signal specific to linearly bound CO (2176 cm⁻¹) only for 10Ni-In₂O₃ and 15Ni-In₂O₃ (Fig. 2b), while pronounced bands of linear, bridged, (2119 cm⁻¹), and three-fold (2066 cm⁻¹) adsorbed CO was detected for the 5Ni-Al₂O₃ reference. Temperature-programmed desorption of carbon monoxide (CO-TPD, Fig. 2c) corroborated that the contribution of nickel to the adsorption of this molecule is significantly inferior when this metal is in contact with In₂O₃. These findings suggest remarkable electronic effects, i.e., metal-support interactions and/or alloying of nickel and indium⁶, implying high dispersion, for the majority of nickel present, and small particle size for nickel unaffected by indium oxide adsorbing CO.

To shed further light on the promoter’s features, catalysts were analyzed by additional methods. Scanning transmission electron microscopy coupled to energy-dispersive X-ray spectroscopy (STEM-EDX, Fig. 3a) revealed highly and almost homogeneously dispersed nickel in fresh lower-content specimens (1 and 5 wt.% Ni) and more agglomerated structures in fresh higher-content specimens (15 and 20 wt.% Ni). Only in 20Ni-In₂O₃, some of the promoters appeared fully segregated from indium oxide. Slight nickel agglomeration is evident for all samples upon use in the reaction. Investigations by high-resolution transmission electron microscopy (HRTEM) of used materials did not visualize any nickel-based phases in 1Ni-In₂O₃ (Fig. 3b), although the magnification was chosen such that, based on the STEM-EDX results, some nickel must have been present within the imaged areas. In the 5 wt.% Ni sample, some amorphous islands can be found on In₂O₃, which might tentatively correspond to nickel-rich structures, due to their lower contrast compared to In₂O₃. In 15Ni-In₂O₃, a similar phase forms a ca. 1-nm thick layer covering many of the imaged In₂O₃ particles, which are additionally accompanied by poorly crystalline nickel-based particles in the 20 wt.% catalyst.

X-ray absorption near-edge structure spectra (XANES, Fig. 3c) of fresh 15 and 20Ni-In₂O₃ closely resemble that of Ni²⁺ in NiO, while some discrepancies are observed for the lower-content solids, which can be explained by higher dispersion and stronger interaction with In₂O₃. In all but the highest nickel content catalyst, cationic nickel completely transformed into metallic species different from those in pure nickel metal upon use in the reaction, strongly suggesting alloying with indium. In 20Ni-In₂O₃, nickel is present in both metallic and oxidative states, likely because some NiO particles do not fully reduce due to their large size³⁲.
High-resolution X-ray diffraction (XRD, Fig. 3d, Supplementary Fig. 4) measured in monochromatic light ($\lambda = 0.49292$ Å) indicated that NiO, present in the fresh materials, reduced indeed forming InNi$_3$ alloys. Their average crystal size is <1, 2.7 ± 0.7, and 7.3 ± 0.9 nm for the catalysts containing 1, 5, and 15 wt.% nickel, respectively, in good agreement with the microscopy analyses. Alloy formation with indium shall be key to anchor the intermetallic phase strongly to the catalyst surface, preventing significant sintering. Only for 15Ni-In$_2$O$_3$, weak reflections specific to metallic nickel nanoparticles (ca. 1.2 nm) were also observed.
detected, confirming the origin of methane production over this catalyst. Since this analysis necessitated catalyst operation directly within the capillary used for diffraction analysis and the consequent application of significantly lower flow rates than in the reactor, some NiO is still detected in the used solids. However, other characterization techniques mirroring the reaction environment more closely exclude that this phase exists upon reaction for these catalysts.

Analysis of the extended X-ray absorption fine structure (EXAFS, Fig. 3e, Supplementary Table 3) shows Ni–Ni and Ni–In bonds in the first coordination shell of nickel. The total number of first neighbors around nickel is 5.9, 6.5, and 8.6 for 1, 5, and 15Ni-In2O3, respectively. Since 12 neighbors are expected for nickel atoms in the bulk metal and the InNi3 alloy, the alloy on all catalysts shall be well dispersed. Moreover, the number of Ni-Ni bonds progressively increases with higher nickel contents, i.e., 1.4, 2.4, and 3.7 for 1, 5, and 15Ni-In2O3, respectively. This is indicative of stronger nickel interaction with indium and suggests a two-dimensional layer-like morphology for the for the catalysts with lower nickel content, whereas particles additionally exist in higher-content samples.

Volumetric CO2 chemisorption enabled to determine a linear decrease of the exposed In2O3 surface at increasing nickel content, consistent with a Stranski–Krastanov film growth (layer formation)38. Indeed, an asymptotic decrease would be expected if nickel species formed agglomerates following a Volmer-Weber film growth39 (Fig. 3f), as previously observed for the Pd-In2O3 system31. Although In2O3 should be fully covered at a nickel content of 12.5 wt.% (Supplementary Table 4), it is likely that inhomogeneous precursor distribution upon impregnation and the large anisotropy of the In2O3 surface prevented the formation of a perfectly uniform layer. Hence, ca. 40% of the In2O3 surface remained still exposed at a nickel content of 15 wt.%.

With a sound understanding of the DI systems, their structure was further investigated by first principles density functional theory. To represent the two-dimensional nickel phase, Ni(111) layers were placed on top of In2O3(111), identified previously and herein as the most abundant indium oxide termination (Supplementary Table 5)18. Based on the stability of nickel atoms with six neighbors each and to commensurate the oxide lattice (Supplementary Table 6), a slightly compressed nickel layer with 36 atoms per unit cell of In2O3 was considered more relevant than uncompressed layers with fewer nickel atoms (25–27). Interestingly, upon relaxation, nickel atoms in this discrete layer became more densely packed and extracted oxygen atoms from the underlying In2O3 (Fig. 4, Supplementary Movie 1). The driving force for this reconstruction can be traced back to the relative bond strengths, since Ni–In affinities are higher with respect to those of In-In and Ni-Ni (Supplementary Table 7), and is in line with other hydrogenation systems for which distinct but pronounced metal-support interactions were described, such as Cu-ZnO, Pt-TiO2, Pt-CeO2, and Pd-In2O3 (Supplementary Figs. 5 and 6, Supplementary Tables 8 and 9). The oxygen atoms extracted emerge to the outermost catalyst surface and are readily stripped as water by the hydrogen present in the reaction environment, which rationalizes the alloying of nickel with indium. Considering these dynamics, the oxygen atoms on the In2O3 termination were removed prior to placing the nickel layer. The most stable structure was found to be Ni36-In2O3−v, which presents the maximal number of oxygen vacancies (v) on the oxide and nickel atoms (12 and 36, respectively, Supplementary Fig. 7). Still, since it cannot be excluded that a small fraction of the oxygen atoms that emerged at the outermost surface is dynamically stored on the InNi3 patches upon CO2 hydrogenation, the effect of residual four oxygen atoms at the most stabilized positions was also explored (Ni36O4-In2O3−v). Finally, since the presence of isolated nickel species can also not be discarded, nickel atoms were alternatively deposited onto In2O3(111), i.e., Ni1−xInxO2, x = 1–4. These Ni1−xInxO2 structures are less stable than the nickel layer but more stable than NiO under a reducing atmosphere such as upon CO2 hydrogenation. For instance, Ni1−xInxO2 is less stable than Ni36-In2O3−v by +0.36 eV per nickel atom. Overall, the catalysts with up to 10 wt.% nickel are better represented by a combination of Ni36-In2O3−v.
InNi$_3$(111), and In$_2$O$_3$(111), while for higher-content materials InNi$_3$(111) and Ni(111) terminations, along with potentially Ni$_2$-In$_2$O$_3$ (vide infra), are additionally significant.

Reaction mechanism and kinetics. To explain the reactivity of the DI catalysts, the adsorption energies ($E_{\text{ads}}$) of reactants on the distinct models were calculated (>200 adsorption calculations, Fig. 5). Ni(111) is associated with a mildly endothermic CO$_2$ adsorption, which can still be overcome (0.22 eV, Fig. 5), and barrierless hydrogen splitting and will readily transform carbon-based adsorbates into methane. In contrast, CO$_2$ adsorption is significantly weakened on clean InNi$_3$(111) (0.74 eV, Fig. 5), while hydrogen splits more exothermically and without an energy barrier. The chemical behavior of all layered nickel structures simulated is very similar to InNi$_3$(111) (Supplementary Fig. 8). Hence, any alloy type shall be virtually inert in converting CO$_2$ into any product on its own, but can provide hydrogen to the active site of In$_2$O$_3$(111) for coverages lower than one monolayer.

InNi$_3$(111), and In$_2$O$_3$(111), terminations, along with potentially Ni$_2$-In$_2$O$_3$ (vide infra), are additionally significant.

Considering the suppressed CO adsorption ability of the catalysts in CO-DRIFTS and CO-TPD analyses, it is conceivable that some oxygen or hydroxide species populate the alloy film during reaction.

When considering low-content samples, the CO$_2$ hydrogenation performance can be seen as the convolution of that of a (multi)layer of InNi$_3$, pure In$_2$O$_3$, and nickel dimers on In$_2$O$_3$. The unpromoted In$_3$O$_5$ ensembles adsorb CO$_2$ and activate H$_2$ heterolytically. Then, hydrides and protons are transferred to CO$_2$ forming methanol. On InNi$_3$(111) and Ni$_{36}$-In$_2$O$_3$ ensembles, CO$_2$ and CO adsorption are either endothermic or weaker than on In$_2$O$_3$, whereas homolytic H$_2$ adsorption is exothermic and CO$_2$ activated at the In$_3$O$_5$ ensemble on clean In$_2$O$_3$(111) shall be hydrogenated both with hydrogen split on the same active site and with hydrogen spilled from nickel layers, whereby sites at the periphery of the patches will be most relevant in the latter process (Supplementary Fig. 9). The fact that methanol selectivity is higher in nickel-poor catalysts than in nickel-rich ones suggests that hydrides and protons generated on In$_2$O$_3$ are still quite strongly utilized, since hydrogen radicals produced on the alloy favor both methanol and CO formation. In contrast to alloyed phases, low-nuclearity nickel clusters at the In$_3$O$_5$ ensemble, in particular Ni$_2$-In$_2$O$_3$, are expected to be highly active in the competing RWGS (Supplementary Figs. 10 and 11), in striking contrast to low-nuclearity palladium clusters anchored to the same ensemble.

To corroborate the DFT findings, kinetic analyses were carried out over all catalysts to experimentally assess the mechanistic origin of the promotional effect (Fig. 6a). The apparent activation energies for both methanol synthesis and the RWGS reaction,
extracted from catalytic tests conducted at variable temperature (Supplementary Fig. 12), are significantly lower already upon incorporation of the smallest nickel amount in comparison to pure indium oxide (from 101 to 83 and from 116 to 92 kJ mol\(^{-1}\), respectively). The smaller difference between activation barriers for methanol and CO synthesis over 1Ni-In\(_2\)O\(_3\) rationalizes its lower methanol selectivity compared to the unpromoted oxide (from 15 to 9 kJ mol\(^{-1}\)). A further increase of the nickel content has little impact on the activation energies, which reach values of 84 and 80 kJ mol\(^{-1}\) for the RWGS and methanol synthesis for 20Ni-In\(_2\)O\(_3\) respectively. The activation barriers for methanation on the materials active for this reaction (15 and 20Ni-In\(_2\)O\(_3\)) were determined at ca. 90 kJ mol\(^{-1}\), in good agreement with literature on pure nickel catalysts (85–94 kJ mol\(^{-1}\))\(^{35,41,42}\) and corroborating the feasibility of methane formation on metallic nickel sites in these systems. Based on the weak dependence of the apparent activation energies for methanol and CO formation on the nickel content, the mechanisms leading to these products are likely highly similar throughout the materials. Hence, the progressive decay of methanol selectivity in favor of CO at higher loadings (Fig. 1b) has a kinetic origin. Since DFT calculations pointed to facilitated hydrogen activation as the origin of the promotional effect, apparent reaction orders respective to this reactant were determined from experiments at variable H\(_2\) concentrations in the feed (Fig. 6b, Supplementary Fig. 13). For methanol synthesis cannot be enhanced beyond a certain threshold even if more homolytically split hydrogen is provided by more abundant species on-site splitting at working conditions. For the RWGS reaction, the reaction orders increased from −0.7 to −0.3 for 1 and 5Ni-In\(_2\)O\(_3\), i.e., the systems which contain mainly flat InNi\(_3\) structures. On these catalysts, both methanol and CO are produced on unpromoted In\(_2\)O\(_3\) ensembles with the two paths competing for H\(^*\) donated by the alloy patches. The negative reaction orders for the RWGS are explained by its first step (CO\(_2\) + H\(^*\)\,→\,COOH) being kinetically unfavored compared to the first of methanol production (CO\(_2\) + H\(^*\)\,→\,HCO\(_2\))\(^{30}\) (Supplementary Fig. 11a). The progressive, decrease in apparent activation energy for the RWGS reaction upon increasing Ni content might be explained based on the formation of additional metastable Ni\(_1\)-In\(_2\)O\(_3\) and Ni\(_2\)-In\(_2\)O\(_3\) ensembles selective to this competitive reaction (Supplementary Figs. 10 and 11). On Ni\(_x\)-In\(_2\)O\(_3\) ensembles, H\(_2\) splitting is barrierless, and thus the net reaction is not controlled by the partial pressure of H\(_2\). At Ni contents of 15–20 wt.% the RWGS reaction shall be mostly performed on these ensembles rather than on those free of nickel, and the reaction order with respect to H\(_2\) decreases to zero. Overall, the kinetic data are in good agreement with earlier and above-presented investigations. Specifically, when H\(_2\) is activated on pure In\(_2\)O\(_3\), it is split into polarized species (H\(^{\delta-}\) and H\(^{\delta+}\)) which are adsorbed on the In\(_3\)O\(_5\) ensemble on an In\(_3\)\(^{4+}\) substructure and O\(^{\delta-}\) atom respectively\(^{18}\). The subsequent transfer to adsorbed CO\(_2\) is energetically disfavored due to the strong polar interaction between adsorbed H\(_2\) and the In\(_2\)O\(_3\) surface. Neutral hydrogen atoms, provided by the alloy phases, do not have to overcome this energy barrier, thus leading to lower activation energies for methanol synthesis and the RWGS reaction\(^{32}\). However, consecutive proton and hydride supply to adsorbed CO\(_2\) are highly selective towards methanol, whereas uncharged species foster both methanol and CO formation. Consequently, methanol synthesis cannot be enhanced beyond a certain threshold even if more homolytically split hydrogen is provided by more abundant alloy phases. On the contrary, the RWGS reaction is boosted to a greater extent in the presence of high nickel contents, presumably as a consequence of an excessive supply of hydrogen atoms (Fig. 6c).

Fig. 6 Nanostructure-driven kinetics and selectivity of nickel-promoted In\(_2\)O\(_3\) catalysts. a Apparent activation energies (\(E_c\)) and b reaction orders respective to H\(_2\) (\(n_{H_2}\)) for methanol synthesis and the RWGS reaction over In\(_2\)O\(_3\) catalysts as a function of their nickel content. c Graphical representation of the structures of In\(_2\)O\(_3\) and Ni-In\(_2\)O\(_3\) catalysts with low and high nickel loadings and indications where products shall be predominantly formed on them. The rings surrounding the molecules represent the respective product selectivity. Reaction conditions: \(T = 553\) K, \(P = 5\) MPa, WHSV = 24,000 cm\(^3\)STP\,g\(^{-1}\)cat\,h\(^{-1}\).
Discussion

Herein, the use of nickel as an economically attractive promoter for In$_2$O$_3$ in the direct hydrogenation of CO$_2$ to methanol was studied in fundamental and applied terms. Considering facile synthetic strategies, dry impregnation led to more stable and active catalysts than coprecipitation. Methanol synthesis was boosted along with the RWGS reaction to some extent and no methane was formed below a nickel content of 10 wt.%, despite the known high CO$_2$ methanation activity of nickel nanoparticles. In-depth characterization revealed a two-dimensional InNi$_3$ phase highly dispersed on In$_2$O$_3$ in nickel-lean samples, which is accompanied by nanoparticles of the same alloy as well as metallic nickel at progressively higher promoter contents. The formation of layered structures rather than agglomerated particles, which is accompanied by nanoparticles of the same alloy as well as metallic nickel at progressively higher promoter contents. The formation of layered structures rather than agglomerated particles, due to peculiar wetting properties of nickel on In$_2$O$_3$ fostering film growth, and their strong anchoring on the oxide via alloying emerged as key contributors to the high catalyst stability.

DFT simulations elucidated that indium-modulated nickel layers easily provide homolytically split hydrogen to In$_2$O$_3$, enhancing oxygen vacancy formation and contributing to CO$_2$ hydrogenation, while barely activating CO$_2$ on their own, which overall explains the beneficial effects and the lack of methane generation. Hydrogen radicals spilled from the InNi$_3$ phase can comitantly support methanol and CO formation, while hydrides and protons produced on In$_2$O$_3$ preferably mediate methanol production. The variable relevance of the former species at distinct contents rationalizes the product distribution and kinetic parameters experimentally determined across all samples. The catalyst comprising 1 wt.% of nickel offers an optimal balance between charged and radical hydrogen atoms, reaching a doubled methanol STY compared to unpromoted indium oxide. Overall, this study identified key structural and electronic features controlling the performance of the classical hydrogenation metal nickel in contact with indium oxide relevant to attain a stable promoted system for a sustainable application. It also highlights that the atomic engineering of a promoter for In$_2$O$_3$ is strongly metal-specific, even when similarity in behavior is expected for elements belonging to the same group in the periodic table.

Methods

Catalyst preparation. Unpromoted and nickel-promoted (1−2.5 wt.% nickel) In$_2$O$_3$ catalysts were prepared via a (co)precipitation (CP) synthesis similar to one reported earlier.

In addition, 1−20 wt.% and 5 wt.% nickel was added to pure indium oxide and mixed indium-aluminum oxide supports with variable stoichiometry (0−100 mol% indium), respectively, by a dry impregnation (DI) method. The nickel-containing catalysts are labeled with the amount of promoter in wt.% separated by a hyphen from the carrier, i.e., Sn-In$_2$O$_3$ indicates 5 wt.% nickel on In$_2$O$_3$. Details to all syntheses applied are provided in the Supplementary Methods.

Catalyst characterization. The metal content in the catalysts was determined by XRF, and porous properties of the catalysts were assessed by N$_2$ sorption. Catalyst reducibility was monitored by temperature-programmed reduction in hydrogen (H$_2$-TPR). CO adsorption was assessed by diffuse-reflection Fourier transform infrared spectroscopy (CO-DRIFTS) and temperature-programmed desorption (CO-TPD). Nickel speciation, coordination geometry, and dispersion were accessed via X-ray absorption spectroscopy (XAS), X-ray diffraction in monochromatic light (XRD), STEM-EDX, and HRTEM. The surface area of In$_2$O$_3$ accessible to reactants was determined using volumetric chemisorption of CO$_2$. Details to all characterization techniques are available in the Supplementary Methods.

Catalytic evaluation. The experimental setup used for catalytic testing is described in detail elsewhere.

Briefly, all experiments were performed in a high-pressure continuous-flow fixed-bed reactor with an inner diameter of 2.1 mm surrounded by an electric furnace. In a typical experiment the reactor was loaded with 100 mg of catalyst with a particle size of 75−100 μm, which was held in place by a bed of quartz wool and heated from ambient temperature to 553 K (5 K min$^{-1}$) at 5 MPa under a flow of H$_2$ (20 cm$^3$ STP min$^{-1}$). After 3 h, the gas flow was switched to the reactants mixture (40 cm$^3$ STP min$^{-1}$) comprising H$_2$ and CO$_2$ in a molar ratio of 4:1. To determine apparent activation energies, the reaction was initiated at 473 K and the temperature stepwise increased to 553 K (increments of 20 K). Reaction orders with respect to H$_2$ were acquired applying a constant flow of CO$_2$ (6 cm$^3$ STP min$^{-1}$) and increasing the flow of H$_2$ (from 20−32 cm$^3$ STP min$^{-1}$, increments of 3 cm$^3$ STP min$^{-1}$), while using He to balance the total flow to 40 cm$^3$ STP min$^{-1}$. Ethane (0.5 cm$^3$ STP min$^{-1}$, Messer, >99.9%) was added to the effluent stream to serve as an internal standard before the stream was sampled every 20 min and analyzed by online gas chromatography. The evaluation procedure of gas chromatography data is reported in the Supplementary Methods. Materials were tested for 16 h for performance comparison, catalyst stability was established over 72 h on stream, and, during kinetic tests, data were collected for 3 h at each condition and averaged. The findings of this study are available within the article and its Supplementary Information file.

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References

1. van Santen, R. A. Chemical basis of metal catalyst promotion. Surf. Sci. 251–252, 6–11 (1991).

2. Bartholomew, C. H. & Farrauto, R. J. Fundamentals of Industrial Catalytic Processes. (Wiley, Hoboken, 2006).

3. Hutchings, G. J. Promotion in heterogeneous catalysis: A topic requiring a new approach? Catal. Lett. 75, 1–12 (2001).

4. Zeraa, F. Shape-controlled nanostructures in heterogeneous catalysis. ChemSusChem 6, 1797–1820 (2013).

5. Pérez-Ramírez, J. & López, N. Strategies to break linear scaling relationships. Nat. Catal. 2, 971–976 (2019).

6. Dong, C. et al. Supported metal clusters: Fabrication and application in heterogeneous catalysis. ACS Catal. 10, 11011–11045 (2020).

7. Zhang, Z., Zandkarimi, B. & Alexandrova, A. N. Ensembles of metastable states govern heterogeneous catalysis on dynamic interfaces. Acc. Chem. Res. 53, 447–458 (2020).

8. Kondratenko, E. V., Mul, G., Baltrusaitis, J., Larrazábal, G. O. & Pérez-Ramírez, J. Status and perspectives of CO$_2$ conversion into fuels and chemicals by catalytic, photocatalytic and electrocatalytic processes. Energy Environ. Sci. 6, 3112–3135 (2013).

Data availability

The authors declare that the data supporting the findings of this study are available within the article and its Supplementary Information file. The DFT data are accessible at the ioChem-BD database at https://doi.org/10.19061/iochem-bd-1-183. All other relevant source data are available from the corresponding author upon reasonable request.

ARTICLE NATURE COMMUNICATIONS | https://doi.org/10.1038/s41467-021-22224-x | www.nature.com/naturecommunications
9. Álvarez, A. et al. Challenges in the greener production of formates/formic acid, methanol, and DME by heterogeneously catalyzed CO₂ hydrogenation processes. *Energy Environ. Sci.* 12, 3425–3436 (2019).

11. Dias, V. et al. Energy and economic costs of chemical storage. *Front. Mech. Eng.* 6, 21 (2020).

12. Martin, O. et al. Indium oxide as a superior catalyst for methanol synthesis by CO₂ hydrogenation. *Angew. Chem., Int. Ed.* 55, 6261–6265 (2016).

13. Frei, M. S. et al. Role of zirconia in indium oxide-catalyzed CO₂ hydrogenation to methanol. *ACS Catal.* 10, 1133–1145 (2020).

14. Tsoukalou, A. et al. Operating X-ray absorption spectroscopy identifies a monoclinic ZrO₂Sn solid solution as the active phase for the hydrogenation of CO₂ to methanol. *ACS Catal.* 10, 10060–10067 (2020).

15. Alibani, D. et al. Semi-hydrogenation of acetylene on indium oxide: Proposed single-ensemble catalysis. *Angew. Chem., Int. Ed.* 56, 10755–10760 (2017).

16. Zhang, S. et al. Solid frustrated-lewis-pair catalysts constructed by regulations on surface defects of porous nanorods of CO₂. *Nat. Commun.* 8, 15266 (2017).

17. Wang, L. et al. In₂O₃ nanocrystals for CO₂ hydrogenation to methanol. *J. Catal.* 361, 313–321 (2018).

18. Wang, L. et al. In₂O₃ nanocrystals for CO₂ fixation: atomic-level insight into the role of grain boundaries. *Science* 360, 390–398 (2019).

19. Ye, J., Ge, Q. & Liu, C.-J. Effect of PdIn bimetallic particle formation on CO₂ reduction over the Pd-In/In₂O₃ catalyst. *Chem. Eng. Sci.* 135, 193–201 (2015).

20. Rui, N. et al. CO₂ hydrogenation to methanol over Pd/In₂O₃: Effects of Pd and oxygen vacancy. *Appl. Catal., B* 218, 488–497 (2017).

21. Snider, J. L. et al. Revealing the synergy between oxide and alloy phases on the role of grain boundaries. *Angew. Chem., Int. Ed.* 58, 9904–9905 (2019).

22. González-Garay, A. et al. Plant-to-planet analysis of CO₂-based methanol production. *Green Chem.* 21, 3370–3378 (2019).

23. Bergwerff, J. A. & Weckhuysen, B. M. In *Handbook of Heterogeneous Catalysis* (eds. Ertl, G. et al.) 1188–1192 (Wiley-VCH, 2013).

24. Hengstebeck, H. et al. Hydrogenation of CO₂ to methanol on a Au-promoted indium oxide catalyst without coke formation. *J. Catal.* 177, 460–472 (1998).

25. Perdew, J. P., Burke, K. & Ernzerhof, M. Generalized gradient approximation made simple. *Phys. Rev. Lett.* 77, 3865–3868 (1996).

26. Blöchl, P. E. Projector augmented-wave method. *Phys. Rev. B* 50, 17953–17979 (1994).

27. Kresse, G. & Furthmüller, J. Efficiency of ab initio total energy calculations for metals and semiconductors using a plane-wave basis set. *Comput. Mater. Sci.* 6, 15–50 (1996).

28. Weatherbee, G. D. & Bartholomew, C. H. Hydrogenation of CO₂ on group VIII metals: II. Kinetics and mechanism of CO₂ hydrogenation on nickel. *J. Catal.* 177, 460–472 (1998).

29. Kresse, G. & Furthmüller, J. Efficient iterative schemes for ab initio total-energy calculations using a plane-wave basis set. *Phys. Rev. B* 54, 11169–11186 (1996).

30. Martin, O. et al. Hydrogenation of CO₂ to methanol over In₂O₃ catalyst. *J. Catal.* 121, 391–402 (1993).

31. Weatherbee, G. D. & Bartholomew, C. H. Hydrogenation of CO₂ on group VIII metals: I. Kinetics and mechanism of CO₂ hydrogenation on nickel. *J. Catal.* 177, 460–472 (1998).

32. Henkelman, G., Uberuaga, B. P. & Jónsson, H. A climbing image nudged elastic band method for finding saddle points and minimum energy paths. *J. Chem. Phys.* 113, 9901–9904 (2000).

33. Heyden, A., Bell, A. T. & Keil, F. J. Efficient methods for finding transition states in chemical reactions: comparison of improved dimer method and partitioned rational function optimization method. *J. Chem. Phys.* 123, 224101 (2005).