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The engineering of CO₂ hydrogenation catalysts for higher alcohol synthesis

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Anthropogenic CO₂ emissions have drawn significant attention in recent years. Using CO₂ as feedstock for chemical processes has become a key solution in overall closed carbon cycles for a vision of a circular carbon economy. CO₂ hydrogenation to higher alcohols has emerged as one of the most promising CO₂ conversion pathways for mitigating CO₂ emissions and producing value-added chemicals. The present review critically discusses the most recent cutting-edge catalyst development in higher alcohol synthesis (HAS), focusing on the influence of different metals, promoters, and supports according to the contributions of different active species in modern catalyst configurations. Particularly, the critical roles of oxygen vacancies and the reaction mechanisms shed light on the rational design of the next-generation CO₂ hydrogenation catalysts.

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Sustainability spotlight

Utilizing CO₂ as feedstock integrated with renewable H₂ is one of the most promising pathways for a closed carbon loop cycle for the circular carbon economy. This review provides a critical assessment on the most emerging CO₂ hydrogenation catalyst designs, discussing the importance of appropriate combination of metal active sites, supports, and promoters for tailoring metal interfaces, metal support interactions, and acid–base sites, *etc.* The review also delves into the reaction pathways that balance between dissociative and nondissociative CO activation, and promote CO insertion and C–C coupling for higher alcohol formation. We believe the readers will find the insights in this review paper enlightening and useful for tailoring advanced catalysts with precise design.

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Introduction

Excessive carbon dioxide (CO₂) emissions, one of the primary greenhouse gases, have caused serious environmental concerns, which result in global warming and climate change.



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The atmospheric CO₂ concentration has continued to rise over the past decades, exceeding 410 ppm.¹ Establishing a practical framework for mitigating CO₂ emissions has been periodically discussed at the governmental level. Most nations have signed the Paris Agreement during the UN Climate Change Conference (COP21), which emphasizes limiting global warming to well below 2 degrees Celsius by restricting cumulative CO₂ emissions to less than one trillion tons of CO₂.² According to recent climate reports, the global average temperature is projected to rise by over 2 °C by 2050 and more than 4 °C by 2100.³ Considerable attention has been directed towards stabilizing atmospheric CO₂ levels, not only by reducing CO₂ emissions but also by effectively utilizing CO₂. CO₂, as the main industrial C₁ carbon source, can be utilized in a wide range of applications.

Over the past few decades, there has been a significant rise in research studies on CO₂ hydrogenation, demonstrating the effectiveness of CO₂ hydrogenation reactions in reducing CO₂ emissions. CO₂ hydrogenation of valuable chemicals has been widely explored using catalytic CO₂ conversion technologies such as thermocatalytic conversion, photocatalytic conversion, and electrocatalytic conversion.⁴ Thermocatalytic conversion, among these technologies, is the most promising due to its high efficiency and great opportunities in large-scale applications.^{5,6} Thermocatalytic CO₂ conversion also promotes a more sustainable approach by utilizing renewable green hydrogen generated *via* wind power, photovoltaic cells, or excess nuclear power from water electrolysis systems.^{7–9}

CO₂ hydrogenation is an excellent strategy that mitigates CO₂ release from chemical processes and produces valuable chemicals and fuels, including methane, methanol, ethanol, C₂–C₄ alcohols, *etc.*, depending on the nature of the specific catalytic active sites and the reaction conditions.¹⁰ Particularly, the CO₂-to-methanol route has been a promising industrial process due to the significant demand for methanol.¹¹ On the other hand, higher alcohols (C₂–C₄ alcohols) offer benefits over

methanol, including greater energy density,¹² lower azeotropic behaviour,¹³ and less water affinity,¹² making them strong alternatives for methanol substitutes in fuels and fuel additives. Also, synthesizing higher alcohols, especially ethanol, at low temperatures is thermodynamically more desirable than methanol due to lower Gibbs free energy and a greater equilibrium constant value.¹⁴ Additionally, higher alcohols are used as solvents for resins, fats, waxes, ethers, and gums, as well as useful feedstock and intermediates for various chemicals and pharmaceuticals.¹⁵

Despite these benefits, achieving high conversion of CO₂ and high selectivity of higher alcohols has remained challenging. An advanced catalyst design is required to promote carbon chain growth and overcome the high kinetic energy barrier associated with CO insertion through C–C coupling into the carbon chain, ultimately facilitating the formation of ethanol and higher alcohols.^{16,17} To increase CO₂ conversion and selectivity for higher alcohols, it is essential to modify the metal active sites, the metal–metal interfaces, the metal–support interactions, and the physical and chemical characteristics of the catalysts to promote key reaction intermediates through balancing, suppressing or synchronizing reactions including reverse water–gas shift (RWGS), Fischer–Tropsch (FT) synthesis, CO₂ hydrogenation to alkanes and alkenes, methanol synthesis, and higher alcohol synthesis (HAS).

There have been critical reviews regarding CO₂ hydrogenation for HAS.^{14,18,19} Cui *et al.*¹⁸ centered on the various catalytic systems and reaction mechanisms involved by the thermodynamic and kinetic analyses in the conversion of CO₂ into alcohols with discussion focusing on the structure–activity relationship, which was relatively comparable to Latsiou *et al.*¹⁹ but the latter focused solely on HAS. In contrast, Xu *et al.*¹⁴ focused on advancements in catalyst design and performance by categorizing them into different catalyst families with a brief discussion on the promoter and support effect and some



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information on reaction mechanisms. This review aims to provide a comprehensive and systematic analysis of the critical knowledge and research gap that impacts the rational design of effective catalysts for CO₂ hydrogenation to higher alcohols. By summarizing the recent research advancements, we present an integrated and in-depth investigation into the roles of different types of metals, promoters, and supports in catalyst system development in HAS through extensive case-by-case studies. We attempt to address this from the viewpoint of stabilizing active phases and structure–performance relationships to improve the production of higher alcohols, hoping that this review may provide insights on the precise engineering of the interplay between active metal sites, supports and promoters at an atomic scale for optimizing HAS catalytic performance. In addition, this review also underlines the influence of oxygen vacancies and emphasizes the reaction mechanism based on *in situ* characterizations and theoretical calculations. Finally, the conclusion and future perspective are explored for future investigation in HAS.

Catalyst design and development

Research on CO₂ hydrogenation for HAS can be categorized into four main groups: (1) modified Cu-based methanol synthesis catalysts,^{20,21} (2) modified Co- or Fe-based FT synthesis catalysts,^{22–27} (3) noble metal-based (Rh, Pd, Ir, Au, and Pt) catalysts,^{28–32} and (4) Mo-based catalyst,^{33,34} as listed in Table 1. Cu-based catalysts are well-known candidates in methanol synthesis and RWGS due to their superior performance and cost-effectiveness.^{34–56} It is now widely recognized that metallic Cu promotes non-dissociative CO activation and its insertion step, thus producing mainly CO and methanol instead of higher alcohols. Modifications of Cu-based catalysts have been developed to enhance the selectivity of higher alcohols.¹⁸ Highly dispersed Cu nanoparticles embedded in the Na-beta zeolite (Cu@Na-beta) were synthesized and tested for CO₂ hydrogenation reaction in a fixed bed reactor.³⁵ The Cu@Na-beta catalyst showed an excellent ethanol selectivity of 79% at 300 °C and 2.1 MPa and a space-time yield (STY) of 398 mg g_{cat}⁻¹ h⁻¹ at a CO₂ conversion of approximately 18%. The authors proposed that the Na-beta can result in the confinement effect of Cu nanoparticles and restrict the formation of unwanted byproducts. Electropositive Cu species were detected in small amounts in the complex Cu@Na-beta catalyst using X-ray photoelectron spectroscopy (XPS) and Cu LMM auger spectra analysis, but none were found in the other prepared catalysts, confirming the cooperative interaction between Cu and Na-beta. The high catalytic performance can be attributed to the synergistic interaction between Cu nanoparticles and the zeolite framework. An *et al.*²¹ reported that the combination of cooperative Cu^I sites on a Zr₁₂ cluster with a metal–organic framework (MOF) exhibited a high ethanol selectivity of >99% with the addition of alkali metal as promoters. Cu^I species are identified as the active sites for ethanol formation, and the role of intimate Cu^I–Cu^I dual sites stabilized by MOF is crucial for inducing the H₂ activation in ethanol synthesis, resulting in the formation of (Cu²⁺–H⁻)² followed by promoting the C–C

coupling of methanol and formyl species. Meanwhile, alkali cation (Cs⁺) creates an electron-rich environment for the Cu center to stabilize the formyl intermediates. This results in the formation of CH₃CHO, which can undergo reductive elimination, producing methanol, ethanol, and water and regenerating the Cu^I–Cu^I sites.

Co-based catalysts have been extensively developed as FT synthesis catalysts due to their strong hydrogenation ability.⁵⁷ Interestingly, several studies suggested that the co-existence of Co⁰ and CoO_x phases in Co-based catalysts can promote the synthesis of ethanol. For instance, Wang *et al.*²⁷ demonstrated that the co-existing phases of Co⁰ and CoO_x in CoAlO_x catalysts are favored in ethanol synthesis. The authors discovered that the different reduction temperatures significantly affected the catalyst performance. CoAlO_x-600 catalyst, which the catalyst reduced at 600 °C, exhibited an outstanding ethanol selectivity of 92.1% and a space-time yield of 0.444 mmol g_{cat}⁻¹ h⁻¹, achieving the best catalytic performance compared to other CoAlO_x catalysts, which were reduced at a temperature between 300 °C and 650 °C (Fig. 1). Moreover, methanol, *n*-propanol and *n*-butanol were also obtained in this catalyst, further validating the effectiveness of Co-based catalysts in CO₂ hydrogenation to higher alcohols. Using the extended X-ray absorption fine structure (EXAFS) spectroscopy, it was observed that the characteristic peak of metallic Co⁰ showed up after reduction at 600 °C with the co-existence of CoO_x species. This suggests that more hydrogen activation can occur due to the presence of metallic Co⁰, resulting in higher catalytic performance. However, CoAlO_x catalysts reduced at temperatures higher than 600 °C, with more metallic sites and a relatively small amount of CoO_x species, encountered an increase in methanol yield but a decrease in ethanol yield.

Similar findings were reported by Ding *et al.*,⁴² indicating the significance of co-existing Co⁰ and CoO_x species in facilitating ethanol formation. The authors successfully synthesized silicalite-1 (S-1) with abundant silanols supported on Co species with 5 wt% and 15 wt% of Co loadings, while the sample synthesized using the same procedure but without the addition of NaOH in the precursor gel was denoted as S–S. Of interest is the capability of this novel approach to utilize strong metal-support interaction (SMSI), thus forming Si–O–Co chemical bonds that can stabilize the co-existence Co⁰ and CoO species. The co-existence Co⁰ and CoO on Co/S-1 was evidenced using characterization techniques such as X-ray diffraction (XRD), CO adsorption (CO-IR), and quasi *in situ* XPS analysis. The catalytic performance of the 5 wt% and 15 wt% Co loadings supported on S-1 showed similar results (Fig. 2). This is consistent with the comparable amount of exposed active sites, *i.e.*, 38.4 μmol g⁻¹ and 38.2 μmol g⁻¹ for 15Co/S-1 and 5Co/S-1, respectively, calculated from H₂-TPD. Besides, 15Co/S-1 and 5Co/S-1 also showed similar results for ethanol selectivity (26% and 27%) and CO₂ conversion (12.1% and 13.9%). A volcano-shaped relationship between CO₂ conversion and the reduction temperature was concluded, highlighting the importance of an optimal reduction temperature to balance the metallic Co⁰ and CoO sites. CO₂ conversion increased as the reduction temperature rose from 200 °C to 300 °C but decreased at higher





Table 1 Summary of reaction conditions and comparison of HAS over selected catalysts

Catalyst	T (°C)	P (MPa)	CO ₂ /H ₂ ratio	GHSV (mL g _{cat} ⁻¹ h ⁻¹)	X _{CO₂} (%)	S _{CO} (%)	S _{C₂H₆} (%)	S _{MeOH} (%)	S _{HA} ^a (%)	STY _{HA} ^a	Ref.
Cu@Na-beta	300	2.1	1:3	12 000	18.0	21.0	—	—	79.0 (ethanol)	398 mg g _{cat} ⁻¹ h ⁻¹ (ethanol)	35
Na-CuCo-9	330	4.0	1:1	5000	20.1	26.5	53.8	—	26.8 (ROH)	80.8 mg g _{cat} ⁻¹ h ⁻¹ (C ₃ H ₇ OH)	36
4.6 K-CuMgZnFe	320	5.0	1:3	6000	30.4	30.6	52.4	1.3	15.9	69.6 mg g _{cat} ⁻¹ h ⁻¹	20
CuFeZn-0.7PDA	310	4.0	1:3	7200	10.6	43.2	22.0	—	34.8 (ROH)	58.2 mg g _{cat} ⁻¹ h ⁻¹	16
Cs-Cu _{0.8} Fe _{1.0} Zn _{1.0}	330	5.0	1:3	4500	36.6	—	—	—	19.8	1.47 mmol g _{cat} ⁻¹ h ⁻¹	37
Cu/ZnAl/K-CuMgZnFe	320	5.0	1:3	6000	42.3	13.8	67.6	1.3	17.4	2.24 mmol g _{cat} ⁻¹ h ⁻¹	38
Cr (1%)–CuFe	320	4.0	1:3	6000	38.4	14.8	—	—	29.2	104.1 mg g _{cat} ⁻¹ h ⁻¹	39
2% Na–Co/SiO ₂	310	5.0	1:3	6000	53.2	—	—	—	12.9 (ROH)	1.10 mmol g _{cat} ⁻¹ h ⁻¹ (ethanol)	40
CoAlO _x -600	140	4.0	1:3	Tank	—	—	—	—	92.1 (ethanol)	0.44 mmol g _{cat} ⁻¹ h ⁻¹ (ethanol)	27
Co/La ₄ Ga ₂ O ₉	270	3.5	1:3	3000	4.6	15.4	34.6	15.3	34.7 (ethanol)	—	41
5Co/S-1	250	2.0	1:3	4670	13.9	0	58.0	—	27.0 (ethanol)	0.83 mmol g _{cat} ⁻¹ h ⁻¹ (ethanol)	42
2.5K5Co–In ₂ O ₃	380	4.0	1:3	2250	36.6	80.8	—	—	—	3.73 mmol g _{cat} ⁻¹ h ⁻¹	43
2% Na–Fe@C/5% K–CuZnAl	320	5.0	1:3	4500	39.2	9.4	—	—	35.0 (ethanol)	—	25
FeNaS-0.6	320	3.0	1:3	8000	32.0	20.7	—	—	12.8	78.5 mg g _{cat} ⁻¹ h ⁻¹	23
K-0.82–FeIn/Ce–ZrO ₂ -900	300	10.0	1:3	4500	29.6	13.4	53.2	4.7	28.7	—	44
ZnFe ₂ O ₄ /Fe–Zn–Na [#]	320	5.0	1:3	12 000	29.1	21.8	—	—	18.6	137.7 mg g _{cat} ⁻¹ h ⁻¹	45
Ir ₁ –In ₂ O ₃	200	6.0	1:5	Tank	—	—	—	—	99.7 (ethanol)	0.99 mmol g _{cat} ⁻¹ h ⁻¹ (ethanol)	46
Pd ₃ Cu	240	3.0	1:3	3000	9.2	—	—	0.8	99.2 (ethanol)	59.2 mg g _{cat} ⁻¹ h ⁻¹ (ethanol)	47
Pd ₃ Cu NPs/P25	200	3.2	1:3	Tank	—	—	—	—	92.0 (ethanol)	41.50 mmol g _{cat} ⁻¹ h ⁻¹ (ethanol)	48
PdFe-6.9 (37 nm)	300	6.0	1:3	6000	33.3	27.9	—	—	19.1	86.5 mg g _{cat} ⁻¹ h ⁻¹	49
0.3K-1Pd/Fe ₂ O ₃	320	4.0	1:3	6000	30.0	23.0	—	—	13.4	52.2 mg g _{cat} ⁻¹ h ⁻¹	26
0.19Na–Rh@S-1	250	5.0	1:3	6000	10.0	—	—	—	24.0 (ethanol)	72.00 mmol g _{cat} ⁻¹ h ⁻¹ (ethanol)	50
2Rh0.5Fe0.5Na/CeO ₂	250	3.0	1:3	6000	9.8	32.1	15.9	27.3	21.5 (ethanol)	63.5 mmol g _{cat} ⁻¹ h ⁻¹ (ethanol)	51
Na–Rh–FeO _x /ZSM-5	200	1.0	—	3600	7.0	—	—	—	97.0 (ethanol)	1.10 mmol g _{cat} ⁻¹ h ⁻¹ (ethanol)	52
K _{0.2} Rh _{0.2} /βMo ₂ C	150	6.0	1:3	Tank	—	—	—	—	72.1 (ethanol)	0.03 mmol g _{cat} ⁻¹ h ⁻¹ (ethanol)	53

^a Selectivity and space-time yield of a specific product are indicated in parentheses.

reduction temperatures. Notably, the Co species in 5Co/S-1 was fully reduced into metallic Co^0 at 400 °C. The increase in CO_2 conversion can be explained by the increase in metallic Co^0 sites, which enhance the dissociation activation of hydrogen on the surface and facilitate the breaking of the C–O bond. However, at higher reduction temperature, aggregation of metals may occur, leading to catalyst deactivation.²⁵ For the same reason, although 5Co/S-1 exhibited superior catalytic activity during the initial phase of the reaction, but it lost stability over time due to gradual reduction of CoO_x to Co^0 under reaction conditions, causing the aggregation of cobalt nanoparticles. In summary, the so-called protection provided by Si–O–Co bond may be able to postpone the reduction of CoO_x to Co^0 , but it is not capable of completely preventing CoO_x from fully reducing into Co^0 , which ultimately affects the catalyst stability.

Previous studies have shown that Pd-based catalysts are effective for CO_2 hydrogenation to methanol and have recently gained attention in HAS. For instance, Lou *et al.*⁴⁷ reported CO_2 hydrogenation to ethanol over Pd/CeO₂ catalysts. In their study, Pd dimers (Pd_2/CeO_2) and Pd nanoparticles (nano-Pd/CeO₂) were supported on CeO₂ with nanorods morphology. Pd_2/CeO_2 catalyst with the unique geometric structure of Pd dimers exhibited impressive catalytic activity with 9.2% of CO_2 conversion, 99.2% of high ethanol selectivity, and STY of 45.6 $\text{g}_{\text{ethanol}} \text{g}_{\text{Pd}}^{-1} \text{h}^{-1}$ at 240 °C and 3 MPa. The STY of atomically dispersed Pd dimers on the CeO₂ catalyst was 90 times higher than that of nano-Pd/CeO₂ with STY of 0.5 $\text{g}_{\text{ethanol}} \text{g}_{\text{Pd}}^{-1} \text{h}^{-1}$ because CO was the primary product formed over nano-Pd/CeO₂ catalyst. High-angle dark-field scanning transmission electron microscope (HAADF-STEM) images further revealed the absence of Pd clusters and particles on both fresh and used Pd_2/CeO_2 catalysts, hypothesizing that Pd species are atomically and uniformly distributed on the CeO₂ support. To the best of our knowledge, the formation of agglomerated Pd clusters and particles can result in a decrease in higher alcohol synthesis. They conclude that the catalytic performance is sensitive to different Pd nanoparticle sizes, with single-atom catalyst (SAC) of Pd dimers exhibiting the best catalytic performance.

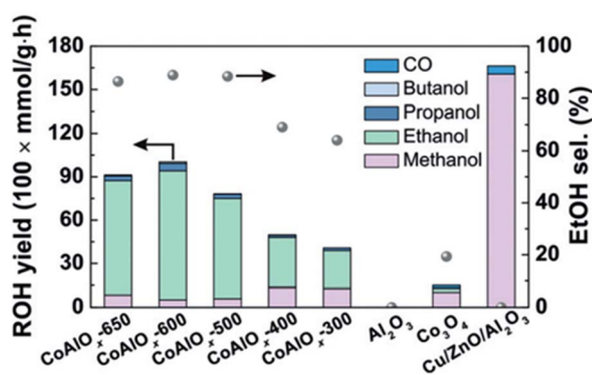


Fig. 1 Catalytic performance of various catalysts based on their alcohols yields and ethanol selectivity. Reprinted with permission from ref. 27. Copyright 2018 John Wiley and Sons, Inc.

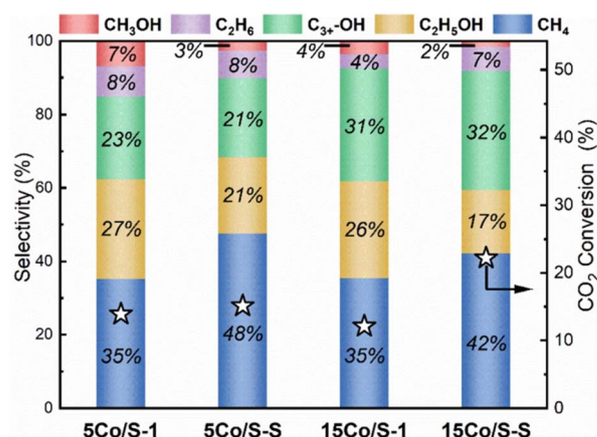


Fig. 2 CO_2 conversion and product selectivity by Co/S-1 and Co/S-S catalysts with different 5 and 15 wt% Co contents. Reprinted with permission from ref. 42. Copyright 2024 Elsevier.

Therefore, this means that tailoring the active sites of a catalyst at an atomic scale can remarkably promote HAS. Meanwhile, the CO adsorption strength, considered the key factor in determining product selectivity, was investigated through *in situ* diffuse reflectance infrared Fourier transform spectroscopy experiments using CO as a probe molecule (CO-DRIFTS). The result demonstrated that the Pd species in Pd_2/CeO_2 catalyst were mostly Pd dimers with no neighbouring Pd atoms, which exhibited stronger CO adsorption strength. Claiming that the stronger binding strength of CO over Pd_2/CeO_2 catalyst can effectively inhibit CO desorption and trigger the C–C coupling reactions between CO and CH_3 intermediate, the formation of ethanol is, therefore, enhanced.

A similar concept has been implemented using an Ir-based catalyst to demonstrate that highly dispersed atomic active sites can promote ethanol generation. In particular, bifunctional Ir₁-In₂O₃ SAC was synthesized through a wet-chemical approach, and the CO_2 hydrogenation reactions were carried out in an aqueous solution.⁴⁶ By impregnating Ir onto the In₂O₃ support, oxygen vacancy (O_v) is induced by partially reducing In₂O₃ in the H₂ atmosphere, as shown in Fig. 3. As a result, the adjacent oxygen vacancy can bind with monoatomic Ir to form Lewis acid–base pair. The partially reduced In₂O₃ and Lewis acid–base pair act as two active sites, which can enhance the CO_2 adsorption and activation into CO^* intermediates. *In situ* DRIFTS experiment results showed further evidence supporting the existence of Ir^{δ+}- CO^* intermediates, while the *in situ* Fourier transform infrared spectroscopy (FTIR) confirmed the existence of $\text{CH}_3\text{O}^*-\text{O}_v$ intermediates. It is concluded that Ir₁-In₂O₃ SAC promotes the C–C coupling by coupling the Ir^{δ+}- CO^* intermediates with the $\text{CH}_3\text{O}^*-\text{O}_v$ intermediates to enhance ethanol selectivity. The CO_2 hydrogenation experiment achieved a high selectivity to ethanol (>99%) and a turnover frequency (TOF) of up to 481 h⁻¹ in this study. The authors also discovered that the ethanol formation was suppressed from 85.3% to 5.7% when the Ir loading increased from 0.2 wt% to 1.0 wt%, which can be ascribed to the agglomeration of Ir nanoparticles and the over-reduction of In₂O₃.



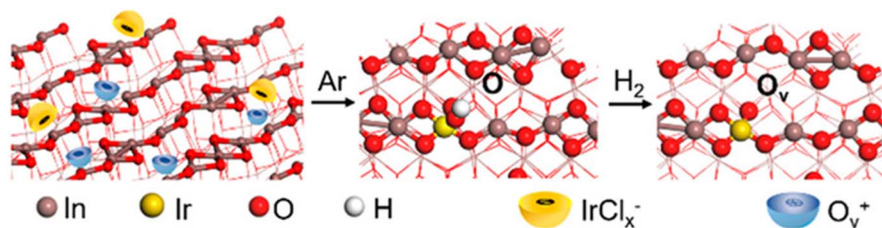


Fig. 3 Schematic illustration of synthesis procedure for Ir₁-In₂O₃ SAC. The diagram depicts the generation of O_v by partially reducing In₂O₃ under H₂ atmosphere prior to impregnating of the Ir. These O_v then act as the anchoring site for single Ir atoms and promote the trapping of IrCl₆²⁻. Reprinted with permission from ref. 46. Copyright 2020 American Chemical Society.

Besides monometallic catalyst systems, bimetallic catalyst systems have recently emerged as promising candidates for CO₂ hydrogenation to higher alcohols. Several studies have shown that introducing a second metal into Cu or noble metal can alter the electronic structure of the catalyst and improve the catalytic performance. Irshad *et al.*³⁶ proposed that in the synthesized Cu-Co bimetallic catalyst, Cu and Co active sites each serve their distinct roles. Na-promoted bimetallic Cu-Co-γ catalysts, combining the capabilities of methanol synthesis and C-C coupling, were prepared with different Cu loadings ($\gamma = \text{Cu}/(\text{Cu} + \text{Co}) \times 10$) *via* coprecipitation method and investigated for the production of higher alcohols. In simpler terms, the Cu site is accountable for the rapid RWGS to produce CO,⁵⁸ and, subsequently, experiencing methanol synthesis. In contrast, Co site is responsible for forming CH_x intermediates through C-O dissociation.⁵⁹ Subsequently, CH₃CO* is formed through C-C coupling between the CO produced and CH_x intermediates, which, after deep hydrogenation, produce acetaldehyde and, ultimately, ethanol or *n*-butanol.^{60,61} Based on their conclusion, acetaldehyde is observed as the key intermediate in the synthesis of *n*-butanol. The “best case” catalyst in their studies was Na-CuCo-9 catalyst, which demonstrated a high C₃₊OH/ROH fraction of 73.5%, giving a high STY_{C₃₊OH} of 80.8 mg g_{cat}⁻¹ h⁻¹ while the overall alcohol selectivity was 58% with CO₂ conversion of 22.1%. Besides, Na-the CuCo-9 catalyst exhibited superior long-term stability without any detrimental effect on its catalytic performance. However, this study did not explore the role of Na as an alkali promoter. The role of alkali metal as a promoter will be further addressed in a later section. On the other hand, Pd-Cu nanoparticles with different composition ratios and supports for ethanol synthesis were investigated in a slurry batch reactor.⁴⁸ By adjusting to the ideal Pd/Cu ratio and catalyst support, the Pd₂Cu/P25 catalyst displayed a high selectivity to ethanol of 92.5% with a TOF value of 359 h⁻¹. The study, however, did not emphasize the formation of the active phase for HAS; instead, it focused on the presence of charge transfer between Pd and Cu. The formation of electronic interaction caused by charge transfer can improve the reduction of surface oxides to their metallic states, which the authors strongly believed to be the active sites for CO₂ activation, thus benefiting the overall CO₂ hydrogenation reaction. Pd₂Cu nanoparticles deposited on various supports such as SiO₂, CeO₂, Al₂O₃, and P25 were also investigated, and the Pd₂Cu supported on the P25 catalyst emerged as the best,

demonstrating the highest TOF and ethanol selectivity out of the four catalysts, which can be attributed to the presence of oxygen vacancies.

Another exciting study is the synthesis of a bimetallic PdFe catalyst.⁴⁹ Iron carbides (FeC_x) are well known as one of the active phases, which positively affects HAS activity, although it is not very active at low CO partial pressures.⁶² Combining FeC_x phases with metal species can form metal-FeC_x interfaces to promote HAS activity. It is presumed that the synergistic effects between PdFe alloy and Fe₅C₂ exist, forming PdFe alloy-Fe₅C₂ interfaces that can eventually promote HAS catalytic performance. Under reaction condition of 300 °C, 5 MPa and WHSV = 6000 mL g_{cat}⁻¹ h⁻¹, higher alcohol selectivity of 26.5% and STY of 86.5 mg g_{cat}⁻¹ h⁻¹ were obtained over PdFe catalyst with Pd loading of 6.9 wt%. The authors demonstrated for the first time that transition metal or alloy could induce the formation of deep iron carbidization without the aid of an alkali ion as a promoter. The presence of PdFe alloy and Fe₅C₂ was verified through *in situ* XRD experiments, showing that Fe₂O₃ in spent PdFe-6.9% catalyst was instantly converted to Fe₃O₄ under hydrogen reduction condition and completely transformed into Fe₅C₂ under CO₂ hydrogenation reaction process within 30 minutes. The XRD analysis also demonstrated that the formation of Fe₅C₂ phase appeared when the Pd loading ≥ 4.3 wt% under CO₂ hydrogenation reaction condition. At the same time, PdFe alloy was formed when reduced Fe atoms from Fe₂O₃ migrated to the Pd surface. The formation of PdFe alloy-Fe₅C₂ interfaces was confirmed through HAADF-STEM images of the spent catalyst. Further evidence of the PdFe alloy-Fe₅C₂ interface formation was shown when the authors compared the HAS activity with the reference catalyst (physically mixing 6.9 Pd/γ-Al₂O₃ + Fe₂O₃ catalyst), which demonstrated no HAS activity. Overall, PdFe alloy mainly generates CO by the RWGS reaction, while Fe₅C₂ is essential for CO dissociation and carbon chain propagation. The adsorbed alkyl is then reacted with CO at the PdFe alloy-Fe₅C₂ interfaces, which is subsequently hydrogenated to ethanol. The authors concluded that PdFe alloy-Fe₅C₂ interfaces are primarily responsible for HAS in their bimetallic system. A study by Yang *et al.*⁴⁵ also observed a similar effect, identifying carburized metal-based catalysts (ZnFe₂O₄/Fe-Zn-Na[#]) as active phases for CO₂ hydrogenation in HAS. Unlike Wang *et al.*⁴⁹ research, in which Fe₅C₂ was reported to be induced after hydrogen reduction and during the CO₂ hydrogenation reaction process, Yang *et al.*⁴⁵ proposed a tandem



catalysis strategy by synthesizing Fe–Zn–Na[#] catalyst (pre-reduced using H₂ for 8 hours) and physically mixing it with ZnFe₂O₄ catalyst to generate FeC_x. However, similar to Wang *et al.*⁴⁹ findings, the Fe₅C₂ phase also appeared after a few hours under reaction conditions as indicated by XRD characterization results. Additionally, CO₂ conversion rapidly increased from 0.3% to 31.8% after 2 h time on stream (TOS) at 300 °C. Higher alcohol selectivity showed a similar pattern, sharply rising from 1.1% to 16.4%. Mössbauer spectroscopy analysis and XRD peaks indicated that the iron species in the spent ZnFe₂O₄/Fe–Zn–Na[#] catalyst were primarily Fe₃O₄, ZnFe₂O₄ and Fe₅C₂. The authors inferred that Fe₃O₄ is responsible for CO formation (RWGS reaction), ZnFe₂O₄ favors the formation of oxygenates, while Fe₅C₂ enhances C–C coupling reaction and initiates CO insertion to produce higher alcohols.

Wang *et al.*²⁵ utilized the multifunctional catalyst of 2% Na–Fe@C/5% K–CuZnAl using a tandem catalysis strategy for CO₂ hydrogenation to ethanol. The Fe@C catalyst was synthesized by pyrolyzing the Fe-based MOFs under a nitrogen atmosphere to promote uniformly distributed Fe active sites due to the periodic arrangement of organic linkers and metallic centers and to control the physicochemical properties of the MOF-derived catalyst. Na doping on a Fe-based catalyst is considered to favour alkene synthesis. Na doping enhances the adsorption of acidic CO₂ due to the formation of a Fe–C bond by electron transfer from Na–Fe@C to CO₂ molecules. Thus, the selectivity of ethanol was slightly enhanced with a slight additional amount of HA produced by adding 2% Na to Fe@C (a 2% Na–Fe@C catalyst). The CuZnAl catalyst was combined with the Na–Fe@C catalyst, which was prepared by mixing two catalytic granules, to boost the synthesis of ethanol from CO₂ hydrogenation. The ethanol selectivity increased from 14% (over a 2% Na–Fe@C catalyst) to 35% (over a 2% Na–Fe@C/5% K–CuZnAl multifunctional catalyst), as shown in Fig. 4. The Na–Fe@C catalyst promotes the reaction of RWGS (over iron oxide, Fe₃O₄) and FTS (over iron carbide, mainly Fe₅C₂), while the addition of the K–CuZnAl catalyst contributes to not only the Fe crystal structure and the catalytic network but also a highly efficient catalyst for the synthesis of methanol to form the intermediate species (formate and CH_xO*).

Furthermore, it is possible that the reaction intermediates produced from one catalytic site could diffuse onto the interface of another catalytic site to synthesize the product over

a multifunctional catalyst during the tandem process. The authors suggested that the reaction intermediates (formate and CH_xO*) could desorb from K–CuZnAl and diffuse onto the interface of the Fe-based catalyst for the possible formation of ethanol through C–C coupling. Moreover, it is necessary to conduct a more in-depth theoretical analysis and *in situ* characterization for multifunctional catalysts due to the highly complex process. Furthermore, the intimacy mode of the 2% Na–Fe@C/5% K–CuZnAl in the cascade reaction produces the highest selectivity and conversion through the granule-mixing mode with 1 g of quartz sand, which may be attributed to active site coverage in the physical-mixing mode of the 2% Na–Fe@C/5% K–CuZnAl. The granule-mixing mode allows for an increase in the spatial distance between distinct catalytic components. The 2% Na–Fe@C/5% K–CuZnAl multifunctional catalyst achieved an ethanol selectivity of 35% with a CO₂ conversion of 39.2%.

Role of promoters or additives

In CO₂ hydrogenation, alkali metals, alkaline earth metals, and transition metals can act as promoters or additives to enhance the overall catalytic activity.⁶³ Some research reported that introducing alkali metals can significantly inhibit hydrocarbon formation by suppressing alkylation reaction, providing opportunities for higher alcohol production.⁴⁴ Some reported that alkali metals could promote structural effects by stabilising active sites or interfaces.⁴⁰ The promotional effect of alkali metals in HAS was intensively studied by different research groups. For example, Zhang *et al.*⁴⁰ synthesized Co/SiO₂ catalysts incorporating alkali metals. Under a CO₂ hydrogenation reaction condition of 250 °C, 5 MPa, and WHSV = 6000 mL g_{cat}⁻¹ h⁻¹, Co/SiO₂ catalyst without any alkali metal promoter and Li–Co/SiO₂ produced only hydrocarbon. In contrast, high selectivity in alcohols, olefins, and CO was achieved over Na–Co/SiO₂ and K–Co/SiO₂ with a decrease in CH₄ selectivity compared to Co/SiO₂ and Li–Co/SiO₂ catalysts. Among the four catalysts, the Na-doped Co/SiO₂ catalyst exhibited the best alcohol selectivity and was further investigated for its role in HAS. It was demonstrated that CO₂ conversion and hydrocarbon selectivity decreased with higher Na contents increasing from 0 to 5 wt% (Fig. 5). The highest alcohol selectivity (12.5%) was obtained over an optimum Na content of 2 wt% Na–Co/SiO₂ catalyst with

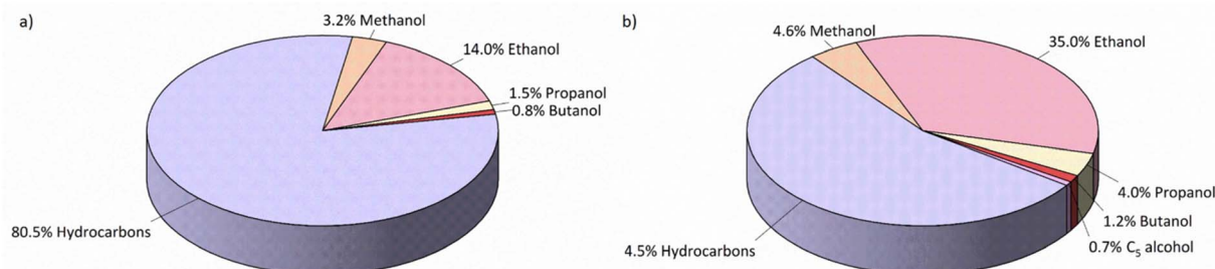


Fig. 4 Product distributions of CO₂ hydrogenation (a) 2% Na–Fe@C catalyst and (b) 2% Na–Fe@C/5% K–CuZnAl multifunctional catalyst (b). Reprinted with permission from ref. 25. Copyright 2021 American Chemical Society.



ethanol STY of $0.47 \text{ mmol g}_{\text{cat}}^{-1} \text{ h}^{-1}$. The most exciting finding in this study was that CO pulse titration experiments showed that introducing Na content improved Co_2C metal dispersion, which aligned with HAADF-STEM and EDX elemental mapping images. Also, a reduction of corresponding particle size was observed. Other observations, such as EXAFS fitting spectra of Co K-edge, further confirmed the improved metal dispersion and reduced particle size as there was a significant decrease in Co–Co coordination numbers in Co_2C from 6.5 to 1.9 when Na content increased from 0 to 5 wt%. All these characterisation results suggested that the Na promoter has a structural effect on Co_2C , which agrees well with other research, highlighting the exceptional ability of alkali metals to exhibit structural effects that stabilize active metal or interface sites.⁶⁴ The authors also concluded that an interaction is formed between Na and Co_2C . This interaction leads to stable Na– Co_2C active sites by generating Na–Co bonds, decreasing particle size, and enhancing the metal dispersion. As a result, RWGS reaction and ethanol STY are improved. Furthermore, an optimized amount of Na is also important in HAS because an excess amount of Na with too strong interaction would weaken the CO adsorption strength and hinder CO insertion, thus leading to high CO selectivity, which is unbeneficial to HAS.

A similar trend was observed in the structure–performance relationship between the Na content and ethanol formation by Zhang *et al.*⁵⁰ They prepared a series of catalysts with different amounts of Na-promoted Rh nanoparticles embedded in zeolite silicalite-1, denoted as $x\text{Na-Rh@S-1}$, where x was the weight loading of Na. By comparing the catalytic performance with different Na content, 0.19Na-Rh@S-1 with a moderate amount of Na content displayed the best catalytic performance in ethanol generation among all, with CO_2 conversion of 10% and ethanol selectivity of 24%. Although the results were generally lower than other published works on HAS, the ethanol STY over 0.19Na-Rh@S-1 was surprisingly high, achieving $72 \text{ mmol g}_{\text{cat}}^{-1} \text{ h}^{-1}$. C_{3+} alcohols were also detected, but the selectivity was not ideal, being less than 5%. Consistent with prior research, an excessive amount of Na can lead to severe deposition of Na species on the surface Rh sites, causing high coverage by Na^+ on the catalyst surface. Meanwhile, the confinement effect by the rigid framework of zeolite S-1 successfully prevents Rh nanoparticles from agglomeration and restricts the sintering of Na-promoted Rh nanoparticles, supported by TEM results. Hence, Na-Rh@S-1 achieved exceptional long-term stability under reaction conditions. The experimental results (XPS and CO_2 chemisorption) also pointed out the role of Na promoter in inducing the Rh^+ generation, which can coexist with Rh^0 species in the catalyst, thus causing a promotional effect in ethanol synthesis. H_2 -TPR profiles observed that the reduction peaks with Na addition became broader and split into two peaks with increasing Na contents. Besides, the reduction peaks also experienced a right shift to a higher temperature with increasing Na loadings. Taken together, this implies that incorporating Na contents can inhibit the complete reduction of Rh species. Besides, the experiment data also showed that the presence of Na as a promoter improved the CO_2 chemisorption performance, which the presence of strong CO_2 adsorption

strength and Rh^+ species can minimize the H_2 adsorption and dissociation ability, thereby suppressing hydrocarbon formation and enhance C–O bond activation and CO insertion for C–C propagation.^{65,66}

Xu *et al.*²⁰ demonstrated that incorporating a moderate amount of K into CuMgZnFe, abbreviated as K-CMZF catalysts, could effectively boost the production of HAS. It was observed that the selectivity of higher alcohols and STY increased when K loading increased from 0.1 to 4.6 wt% but then decreased at K loading of 17.6 wt%. Optimized K-CMZF catalyst with 4.6 wt% K loading displayed the highest CO_2 conversion of 30.4% and higher alcohols STY of $69.6 \text{ mg g}_{\text{cat}}^{-1} \text{ h}^{-1}$. Based on Anderson–Schulz–Flory (ASF) calculations, the value of chain growth probability (α) increased when K loading increased, highlighting that adding K can enhance the chain growth ability of K-CMZF catalysts. The existence of K promoter not only maintains a great balance between non-dissociative and dissociative CO activation but also controls the hydrogenation capacity, thus ensuring the participation of adequate $^*\text{CH}_x$ and $^*\text{CO}$ species in C–C coupling reactions while restricting $^*\text{CH}_x$ hydrogenation termination reaction to promote $^*\text{CH}_x\text{CO}/^*\text{CH}_x\text{CHO}$ hydrogenation to higher alcohols.

Witton *et al.*⁴³ systematically investigated the promoter effect on CO_2 hydrogenation over K–Co promoted In_2O_3 catalysts. Adding K and Co into In_2O_3 significantly boosts the C–C chain growth and CO insertion, thus increasing the higher alcohol selectivity, but had only a minor effect on CO_2 conversion. On the other side, both Co- and K-promoted In_2O_3 catalysts showed a clear difference in product selectivity, wherein the K-promoted In_2O_3 catalyst was highly selective towards CO and CH_3OH synthesis, while the Co-promoted In_2O_3 favored the production of C_{2+} hydrocarbons over higher alcohols. The authors postulated that the addition of K reduces the weak and medium H_2 adsorption, thereby suppressing the hydrogenation of CH_3OH and formation of CH_4 from CO. Contrarily, characterization results revealed a mixture of Co^0 and CoO in the Co species of Co-promoted In_2O_3 . CoO sites facilitate the formation of C_xH_y^* species, which subsequently react with migrating CO^* on the CoO surface to form C_{2+}OH products. However, the faster hydrogenation of C_xH_y^* species over CO insertion owing to weak H_2 adsorption effectively promotes C_{2+} hydrocarbon formation rather than C_{2+} oxygenates formation. To enhance the HAS activity, K–Co promoted In_2O_3 catalysts with varying weight percent of K and Co were synthesized. As a result, K–Co promoted In_2O_3 (2.5 wt% K and 5 wt% Co) demonstrated superior HAS catalytic performance with higher alcohols STY of $169.6 \text{ g kg}_{\text{cat}}^{-1} \text{ h}^{-1}$ and a higher alcohols distribution of 87.4% among the overall alcohols formed. The improved HAS activity can be attributed to the synergistic interaction between K, Co, and In_2O_3 . The introduction of K and Co contributes to generating K–O–Co sites, which reduces weak H_2 adsorption and causes a stronger interaction of H_2 with the catalyst surface, hence enabling the manipulation of higher alcohol selectivity.

Zhang *et al.*³⁹ examined the dependence of C_{2+}OH selectivity on the Cr loading level of the Cr–CuFe catalyst, in which the maximum C_{2+}OH selectivity was obtained at Cr = 1%. In detail, Cr-promoted CuFe-based catalysts with K additive were



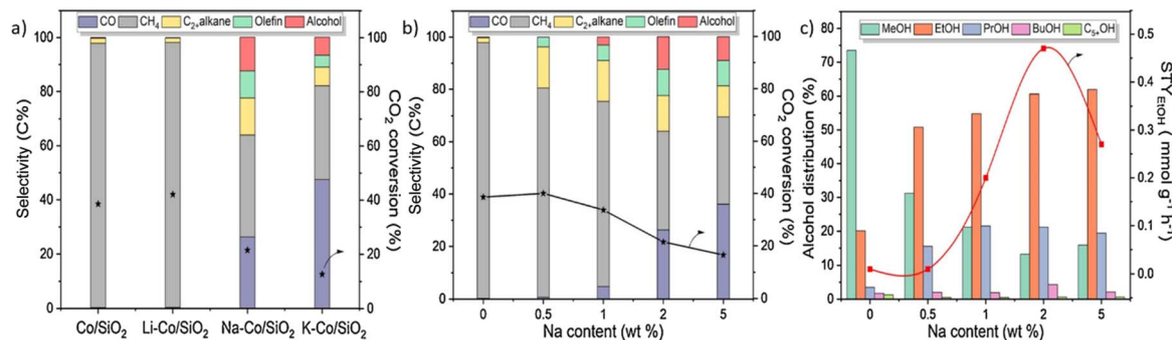


Fig. 5 Catalytic performance over Co/SiO₂ catalysts with different alkali metals and Na contents (a) CO₂ conversion and product selectivity with different alkali metals (b) CO₂ conversion and product selectivity Na contents (c) alcohol distribution and ethanol STY with different Na contents. Reprinted with permission from ref. 40. Copyright 2021 Elsevier.

synthesized *via* the sol-gel method, and the role of Cr as a dopant was further investigated. It was inferred that the interaction between Cu and Fe becomes stronger with the Cr additive. All the CuFe-based catalysts contained the same amount of K as the catalytic performance on Cr (1%)-CuFe catalyst without the K promoter exhibited higher selectivity towards methane and C₂-C₄ alkanes whereas decrease in olefins and higher alcohols selectivity. This is consistent with previous research, indicating the role of alkali metals in preventing over-hydrogenation and hydrocarbon formation. With the aid of alkali metal, catalyst alkalinity is improved, therefore enhancing CO₂ adsorption and activation.²⁰ An appropriate Cr loading amount can enhance H₂ adsorption and activation while weakening CO adsorption strength on the CuFe catalyst. This suggests that Cr additives can effectively mitigate CO over-dissociation, inhibiting the formation of excessive iron carbide species (FeC_x) from metallic Fe. Supported by Fe (2p) XPS and ⁵⁷Fe Mössbauer spectroscopy, FeC_x characteristics were observed over the Cr (1%)-CuFe catalyst compared to the undoped CuFe catalyst. Besides, the amount of Cu⁰ (23.2%) was lower, and the amount of Cu⁺ and Cu²⁺ (76.8%) was higher in Cr (1%)-CuFe catalyst than that of the undoped CuFe catalyst (30.4% of Cu⁰, 69.6% of Cu⁺ and Cu²⁺), as indicated by Cu LMM Auger electron spectra. These findings confirmed the improvement of interactions between Cu and Fe species, which are conducive to forming Cu-FeC_x interfaces, the active sites of the catalyst for HAS.³⁷ Excessive Cr loading has a negative effect on HAS because the Cr species would heavily cover the catalyst surface, and the interaction of CH_x with non-dissociative CO over Cu-FeC_x interfaces will be inhibited. The reaction mechanism proposed that C-C coupling between the interaction between CH_x and non-dissociative CO over Cu-FeC_x interfaces produces more acetate and acetaldehyde intermediates, enhancing the synthesis of higher alcohols.

Ji *et al.*³¹ explored the effects of different transition metals and alkali metals on Rh/CeO₂ catalyst for CO₂ hydrogenation in ethanol synthesis. Fe and Na doping as dual promoters demonstrated superior catalytic activity for CO₂ hydrogenation with high ethanol selectivity of 29.8% and ethanol STY of 116.7 mmol g_{cat}⁻¹ h⁻¹ at 250 °C and 3 MPa. The optimum

amounts of Fe and Na in the RhFeNa/CeO₂ catalyst, denoted as 2Rh0.5Fe0.5Na/CeO₂, were both 0.5 wt%, achieving the highest catalytic performance and Rh dispersions with Rh loading of 2 wt%. H₂-TPR and CO₂-TPD profiles showed that adding Fe could enhance the Rh metal dispersion by showing increased H₂ consumption, while adding Na was also shown to increase H₂ consumption and promote the reduction of Rh species to metallic Rh. Other additives such as Li, Cs, and K suppressed the reduction of Rh species. It was concluded that adding Na and Fe as dual promoters improve the overall Rh metal dispersion and form strong interactions with Rh, thus enhancing the SMSI. FTIR studies of chemisorbed CO and XPS results further revealed that Na and/or Fe additions could enhance the CO adsorption ability, which increased the formation of Rh⁺ active sites. The mechanism studies observed that there were more intermediates of CO* and HCOO* produced on the Fe and Na-promoted Rh/CeO₂ catalysts compared to the unpromoted catalysts, indicating that the doping of Fe and Na promotes dissociation and hydrogenation capacity of CO₂, hence resulting in higher ethanol production, as shown in Fig. 6. However, there was a slight decrease in catalytic activity in the early stage of the reaction, which then became stable after 50 hours. The characterization results of the spent catalyst remained the same as those of the fresh catalyst. The only difference detected was that thermogravimetric analysis (TGA) showed a relatively small weight loss of 6.4% at temperatures ranging between 150 °C and 250 °C, which can be attributed to the desorption of water and organic compound of the spent catalyst surface.

Yao *et al.*²³ also studied the dual promoter effect by incorporating Na and S into monometallic iron catalysts for HAS. The authors proposed that the sulfur existed in the form of sulfate plays a pivotal role in tuning the CO activation through strong synergistic interaction between Na and S, which increases the amount of adsorbed CO to bind with alkyl species on Na-promoted Fe sites, forming higher alcohols. Experimental results in catalytic performance showed that FeNa catalyst exhibited alcohol selectivity of less than 5% while FeNa catalyst promoted with 0.6 wt% of sulfur through precipitation method, denoted as FeNaS-0.6 catalyst, achieved the maximum alcohol



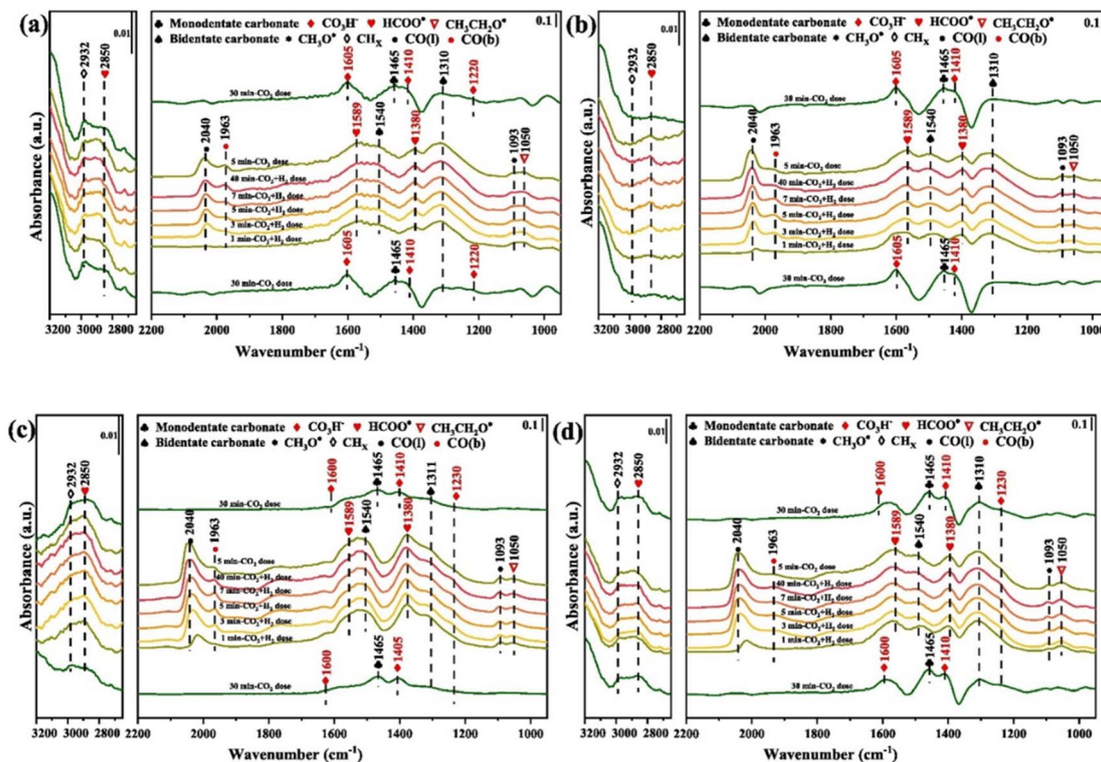


Fig. 6 *In situ* DRIFTS of various catalysts after switching the feed gas from CO₂ to CO₂ + H₂, followed by returning to CO₂ at 250 °C and 0.1 MPa (a) 2Rh/CeO₂ (b) 2Rh0.5Fe/CeO₂ (c) 2Rh0.5Na/CeO₂ and (d) 2Rh0.5Fe0.5Na/CeO₂. Reprinted with permission from ref. 51. Copyright 2024 Elsevier.

selectivity of 16.1% with more than 98% of C₂₊OH fraction in total alcohols. The authors also synthesized the FeNaS-*im* catalyst by introducing the same sulfate content as the FeNaS-0.6 catalyst on FeNa *via* the wet impregnation method to study the promotional effects of sulfate. The higher alcohol selectivity of the FeNaS-*im* catalyst was slightly lower than that of the FeNaS-0.6 catalyst, which can be ascribed to the disadvantage of the impregnation method, resulting in non-uniform sulfate distribution and weak interactions between Fe species and sulfate species. It is well-documented that dissociative and non-dissociative CO activation are essential for HAS. In this study, the presence of sulfate with strong electron-withdrawing ability offers additional Fe sites for non-dissociative CO adsorption. At the same time, adding Na helps maintain the CO dissociative ability of Fe sites, highlighting the importance of maintaining a kinetic balance between dissociative and non-dissociative CO activation in HAS.

Role of catalyst supports

Catalyst support plays a pivotal role in CO₂ hydrogenation by enhancing metal dispersion, stabilizing active sites, regulating catalyst surface interfaces, influencing key intermediates to alter reaction mechanisms, and improving adsorption or activation capabilities.⁶⁷ In addition, the construction of SMSI has been demonstrated to effectively stabilize active sites under CO₂ hydrogenation reaction conditions.^{68,69} The SMSI effect is widely acknowledged for its significant influence on the stability, activity, chemical state, and morphology of the catalysts.⁷⁰ Xin

*et al.*⁶⁸ demonstrated that by constructing an SMSI state, it is possible to promote the formation of a defective MoO_{3-x} overlayer surrounding Ru nanoparticles (Ru@MoO_{3-x}) at a low temperature of 250 °C. This modification favors RWGS, achieving >99% CO selectivity, which could easily revert to CH₄ formation upon O₂ treatment. Therefore, constructing SMSI between active metals and support materials is one of the strategies that could be implemented to increase the higher alcohol selectivity, which also operates under low-temperature reaction conditions. For instance, Zhang *et al.*⁷¹ identified that only SiO₂ and Si₃N₄ supports managed to stabilize the Co₂C active sites by generating Si-O-Co bonds. In their study, the influence of different catalyst supports (Al₂O₃, ZnO, AC, TiO₂, SiO₂, and Si₃N₄) in ethanol synthesis over supported Na-promoted cobalt catalysts was investigated, as illustrated in Fig. 7. Na-Co/Al₂O₃, Na-Co/ZnO, Na-Co/AC and Na-Co/TiO₂ catalysts exhibited very low alcohol selectivity of less than 1%. In comparison, Na-Co/SiO₂ and Na-Co/Si₃N₄ catalysts exhibited a total alcohol selectivity of 9%. The experimental result was explained by the formation of the Co₂C active phase on Na-Co/SiO₂ and Na-Co/Si₃N₄ catalysts, which remained stable even after the reaction. Although the Co₂C active phase was also detected on Na-Co/AC and Na-Co/TiO₂ catalysts, the Co₂C active phase was almost fully decomposed during the reaction condition due to a lack of SMSI. In contrast, no Co₂C active phase was found on Na-Co/Al₂O₃ and Na-Co/ZnO catalysts. The authors inferred that Co₂C is the active phase in CO₂ hydrogenation, with its stability attributed to SMSI facilitated by the



formation of Si–O–Co bonds, which also depends on the nature of the supports utilized.

An *et al.*⁴¹ exploited the SMSI between $\text{La}_4\text{Ga}_2\text{O}_9$ and cobalt particles, facilitating the formation of active $\text{Co}^0\text{--Co}^{2+}$ pairs. Perovskite oxides (PTO), with highly tunable composition and structure, can regulate oxygen vacancies through their bulk and surface properties. $\text{Co}/\text{La}_4\text{Ga}_2\text{O}_9$ catalyst was synthesized using perovskite type $\text{LaCo}_{1-x}\text{Ga}_x\text{O}_3$ support and investigated for CO_2 hydrogenation reaction. Ethanol selectivity of 34.7% was obtained over $\text{Co}/\text{La}_4\text{Ga}_2\text{O}_9$ catalyst under reaction conditions of 270 °C, 3.5 MPa and $\text{GHSV} = 3000 \text{ mL g}_{\text{cat}}^{-1} \text{ h}^{-1}$. Co nanoparticles loaded on $\text{La}_4\text{Ga}_2\text{O}_9$ were formed by reducing the $\text{LaCo}_{0.5}\text{Ga}_{0.5}\text{O}_3$ precursor. SMSI between Co and $\text{La}_4\text{Ga}_2\text{O}_9$ are postulated, resulting in electron donation from cobalt to $\text{La}_4\text{Ga}_2\text{O}_9$, thus forming $\text{Co}^0\text{--Co}^{2+}$ pairs as active sites. CO_2 is adsorbed and activated on the $\text{La}_4\text{Ga}_2\text{O}_9$ surface, which favors RWGS in producing CO intermediates. Then, the produced CO intermediates migrate to $\text{Co}^0\text{--Co}^{2+}$ sites to promote ethanol synthesis. However, it is unfortunate that ethanol yield decreased when reaction time increased. This can be attributed to the increase in the $\text{Co}^0/\text{Co}^{2+}$ ratio, which caused an imbalance in the formation of intermediates over longer periods. Apart from that, SMSI between Co_3O_4 particles and isolated silanols, resulting in the formation of Si–O–Co chemical bonds on $\text{Co}/\text{S-1}$ catalysts, had also been observed by Ding *et al.*⁴² The SMSI was supported by XPS results, which demonstrated an increased number of electronic defects in Co_3O_4 on $\text{Co}/\text{S-1}$ and a higher electron density of Si–O bonds on S-1. This further consolidates the electron transfers from Co_3O_4 to Si–O bonds, contributing to the presence of Si–O–Co chemical bonds. The active sites of the $\text{Co}/\text{S-1}$ catalyst can be tuned through Si–O–Co chemical bonds, which stabilize the co-existing phases of Co^0 and CoO (CoO_x sites) by preventing the complete reduction of Co_3O_4 . DFT calculations revealed the roles of Co^0 and CoO sites in CO_2 hydrogenation to ethanol, respectively. Specifically, Co^0 sites favor the COOH^* species, while CoO sites promote the formation of HCOO^* species. The COOH^* and HCOO^* species are then coupled with CH_x^* species on CoO_x sites to produce ethanol.

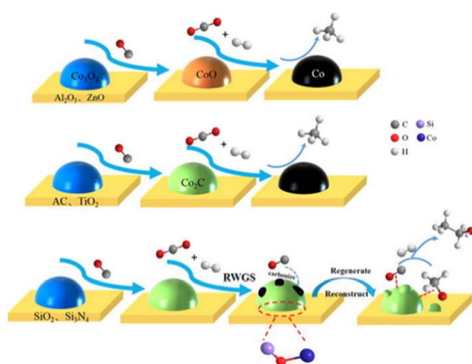


Fig. 7 Schematic of Co phase transformation during CO reduction and CO_2 hydrogenation reaction on Na–Co catalysts supported on different supports. Reprinted with permission from ref. 71. Copyright 2020 Elsevier.

Moreover, a few studies have investigated the confinement effect of supports to achieve catalyst stability in CO_2 hydrogenation. Wang *et al.*⁷² recently postulated that the confinement effect in $\text{In}_2\text{O}_3\text{--TiO}_2$ catalysts promotes the dispersion of the In_2O_3 guest catalyst onto the TiO_2 host surface, thereby improving performance in CO_2 hydrogenation to CO. The formation of In–O–Ti interfacial bonds was crucial in inducing In_2O_3 dispersion and stabilizing the metastable InO_x layers. Their findings revealed that the confinement effect at oxide/oxide interfaces benefits CO_2 hydrogenation. For CO_2 hydrogenation in HAS, Ding *et al.*³⁵ synthesized the $\text{Cu}@/\text{Na-beta}$ catalyst, embedding the Cu nanoparticles in the beta molecular sieve. The authors highlighted that the synergistic interaction between copper nanoparticles and the beta zeolite framework is essential for HAS catalytic performance as this synergy suppresses the CO_2 adsorption site and prevents the over-reduction of the catalyst. Na-beta zeolite as support in $\text{Cu}@/\text{Na-eta}$ catalyst demonstrates a superior confinement effect that shapes the copper nanoparticles into unique configurations with surface sites and limits the reactants along the catalyst surface, thus successfully preventing the formation of by-products such as methanol, formic acid, or acetic acid. In another study, Cu nanoparticles were embedded into BEA and MFI (S-1) zeolites, and their mesoporous equivalents (m-zeolites) were obtained *via* the carbon templating method.⁷³ Based on the catalytic test result, mesoporous zeolites exhibited a higher CO_2 conversion and ethanol selectivity than their non-mesoporous counterparts. Besides, the introduction of mesopores facilitated the formation of isopropanol. Cu-based mesoporous S-1 zeolite ($\text{Cu}@/\text{m-S1}$), which underwent an additional recrystallization procedure, achieved a higher isopropanol selectivity and yielded $20.51 \text{ mmol g}_{\text{cat}}^{-1} \text{ h}^{-1}$. The authors claimed that the mesoporous effect of zeolites positively affects the production of alcohol by facilitating the accessibility of reactant access to Cu active sites and improving the SMSI to accelerate alcohol formation. Tran *et al.*⁵² developed a Na–Rh– $\text{FeO}_x/\text{ZSM-5}$ catalyst, which exhibited a higher ethanol selectivity than a Na–Rh– FeO_x catalyst without zeolite support. The high specific surface area, smaller pore size, and decrease in iron particle size can be attributed to the excellent synergy between the Na–Rh– FeO_x and ZSM-5 support. The ZSM-5-supported catalyst showed a suppression effect on hydrocarbon formation. However, the CO_2 conversion did not increase despite the higher density of active sites due to higher surface area. This indicates that the zeolite support can modify the properties of the active sites.

Critical roles of oxygen vacancies

Various catalytic materials (*e.g.*, Rh, Co, Fe, Cu, and Au) are active sites to catalyze CO_2 hydrogenation and produce higher alcohols.^{16,29,74,75} However, monometallic catalysts have limitations and restrictions for HAS from CO_2 , such as low conversion and selectivity. Thus, metal oxides (*e.g.*, ZrO_2 , TiO_2 , and CeO_2) are often used as supports or promoters due to their inherent ability to generate abundant oxygen vacancies. The oxygen vacancy is typically recognized as the active site for CO_2



adsorption and activation, facilitating the formation of intermediates and promoting SMSI, hence assisting in accelerating CO₂ hydrogenation.^{41,44,76,77} Yang *et al.*¹⁶ synthesized the N-doped CuFeZn catalyst and added an organic ligand, 2,6-pyridine dicarboxylic acid. Accordingly, adding PDA to the CuFeZn catalyst increases the generation of oxygen vacancies, which regulates the electron structure of the catalyst. The oxygen vacancy captures CO₂ by trapping O, and the nearby Zn or Cu reacts with the captured CO₂ to form CO₂^{•-} or CO* species. This is most likely because O in CO₂ recharges the oxygen vacancy and strengthens the interaction between O and the active metals of the catalyst, assisting in establishing the pathway for electron transfer, facilitating the activation, dissociation, and reduction of CO₂ molecules, and thus promoting the formation of higher alcohols. Xi *et al.*⁴⁴ reported the K-promoted Fe-In/Ce-ZrO₂ catalyst in which the existence of reduced In_xO_x phases with oxygen vacancies is of great importance for CO₂ activation; thereby, these oxygen vacancies will assist CO₂ activation in the formation of key reaction intermediates such as HCOO*, H₂COO*, and H₂CO* species. This study demonstrated that *in situ* pretreatment with CO at 350 °C for 3 h boosted the oxygen vacancy by 348.6% compared to air-calcination treatment. Moreover, Li *et al.*¹⁷ reported tailoring the Fermi level of a Cu-doped Zn catalyst synthesized using the urea homogeneous hydrolysis technique, suggesting that a lower Fermi level promotes better carbon chain growth. The electron trap effect of oxygen vacancy regulates the Fermi level of semiconductors. It influences their catalytic performance *via* charge transfer between the surface and adsorbed substances with an imbalanced electron structure, resulting in fast electron transfer. Thus, oxygen vacancies could boost the formation of higher alcohols. Besides, surface oxygen vacancies also play a key role in promoting CO₂ adsorption and activation. In the Ir₁-In₂O₃ SAC catalyst, the oxygen vacancies combine with monoatomic Ir to form Lewis acid–base pairs, which serve as active sites for CO₂ adsorption and activation into CO* intermediates. The

Ir^{δ+}-CO* intermediates interact with the CH₃O*-O_v intermediates, thus enhancing ethanol selectivity.⁴⁶

Reaction mechanisms of CO₂ hydrogenation for HAS

The reaction mechanisms for the synthesis of higher alcohols from CO₂ hydrogenation typically involve the following steps: the initial stage for the activation of CO₂ and H₂, resulting in the formation of C₁ intermediates including CO*, CH_x*, and HCOO* species; and the second stage for the production of ethanol or higher alcohols with a longer carbon chain (C₂₊-C₄₊) through C-C coupling of CH_x* - CO* (a CO-mediated pathway) or CH_x* - HCOO* (a formate-mediated pathway).^{17,29,37,44,71,78} Yang *et al.*¹⁶ proposed reaction mechanisms for HAS from CO₂ hydrogenation over N-doped CuZnFe, as illustrated in Fig. 8. The active sites (*e.g.*, Cu⁰, ZnO, Fe₃O₄, and Fe₅C₂) adsorb the reactant molecules. Through capturing O from CO₂, the oxygen vacancies generated on the surface of ZnO facilitate the adsorption and activation of CO₂ molecules. The weak base of doped N also contributes to activating CO₂ molecules. This leads to the generation of CO₂^{•-} species as CO₂ captures electrons over the surface of ZnO or N. At the interface of Cu-ZnO, H₂ is dissociated and activated due to the vacant s orbital of the Cu species. Through the RWGS reaction, at this interface of Cu-ZnO, CO₂ is dissociated to form CO* species. Subsequently, CO* species may undergo hydrogenation to form methanol or desorb to produce CO. Furthermore, at the Cu-Fe₂C₂ interface, a considerable portion of CO* dissociates, resulting in C* and O* radicals. The C* radical is further hydrogenated to generate the alkyl species (CH_x*), and carbon chain growth occurs through continuously inserting CH_x* to form hydrocarbons (C_nH_m). Subsequently, CO* is inserted into C_nH_m on Cu sites to form higher alcohols (C₂₊).

Zhou *et al.*²⁶ proposed a reaction pathway for HAS from CO₂ hydrogenation over a 0.3K-1Pd/Fe₂O₃ catalyst, as illustrated in Fig. 9. The authors investigated the reaction intermediates (*e.g.*, CO, C₂H₅O*, CH₃CHO*, CH₃CH₂O*, and CH₄) after injecting

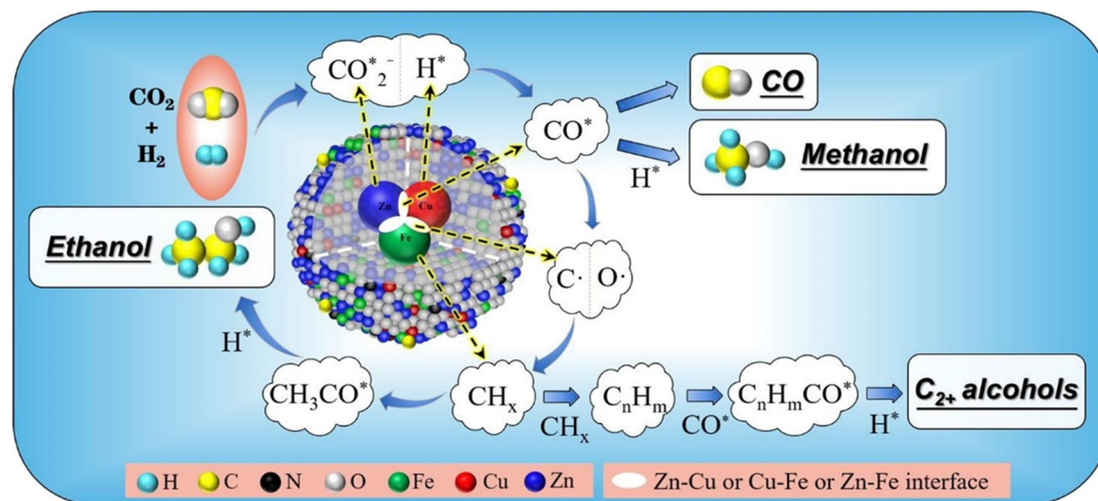


Fig. 8 The weak base of doped N promotes the generation of CO₂^{•-} species through CO₂ activation, which is further regulated by the CuZnFe catalyst to form intermediates including CO*, CH_x, and HCOO* species to form higher alcohols through CO* insertion into C_nH_m. Reprinted with permission from ref. 16. Copyright 2022 Elsevier.



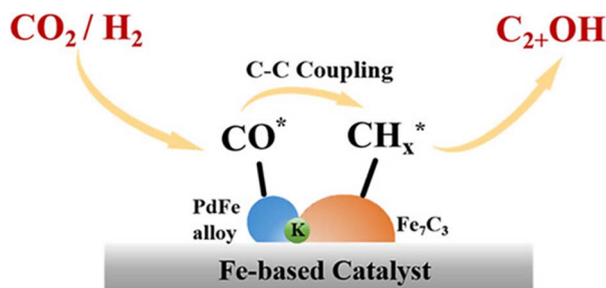


Fig. 9 Proposed reaction pathway that shows the non-dissociative activation of CO regulated by K and Pd to balance the dissociative activation of CO regulated by Fe₂O₃ catalyst for promoting the C–C coupling mechanism. Reprinted with permission from ref. 26. Copyright 2024 American Chemical Society.

CO₂/H₂ over the surface of the catalyst by *in situ* DRIFTS. The introduction of highly dispersed K (0.3 wt%) restricts the hydrogenation of the CH_x* species. It enhances the ratio of CO*/CH_x*, promoting the insertion of CO, while introducing 1% Pd increases the formation rate of C₂H₅* species. The synergy of K and PdFe alloys on a 0.3K–1Pd/Fe₂O₃ catalyst can regulate the ratio of CO*/CH_x* on the surface of the catalyst. The observed key reaction intermediates by *in situ* DRIFTS suggest the reaction pathway following the CO-mediated

reaction mechanism by the insertion of CO* into CH_x* species to form intermediates (CH_xCO*) through C–C coupling, which may instantly conjugate CO* and CH_x* species, resulting in the formation of higher alcohols. Additionally, the study performed CO-TPSR-MS experiments to investigate the influence of K and Pd loading on Fe₂O₃ towards the behavior of the dissociative and non-dissociative activations of CO by measuring the ratio of CO₂/CH₄ from the desorption peak area. The signals CH₄ and CO₂ signify the dissociative activation of CO and the non-dissociative activation of CO, respectively. Fe₂O₃ shows a strong dissociative activation of CO. The addition of 1% Pd to 1Pd/Fe₂O₃ slightly enhances the non-dissociative activation of CO, which is attributed to the formation of PdFe alloys. However, adding 0.3% K to 0.3K/Fe₂O₃ shows a significant non-dissociative activation of CO. The CO-TPSR-MS experiments indicate that the synergy of K and PdFe alloys along with Fe₇C₃ regulates a favorable ratio of CO*/CH_x* on the surface of the 0.3K–1Pd/Fe₂O₃ by balancing the dissociative and non-dissociative activations of CO. This facilitates the C–C coupling of CO* and CH_x* species to produce higher alcohols.

Xu *et al.*³⁷ proposed an integrated reaction pathway of CO₂ hydrogenation over a Cs–CuFeZn catalyst with a favorable balance of synergistic sites (Cu–ZnO and copper–iron carbide interfaces), as illustrated in Fig. 10. At the CuZnO interfaces, methanol formation proceeds directly from CO₂ through

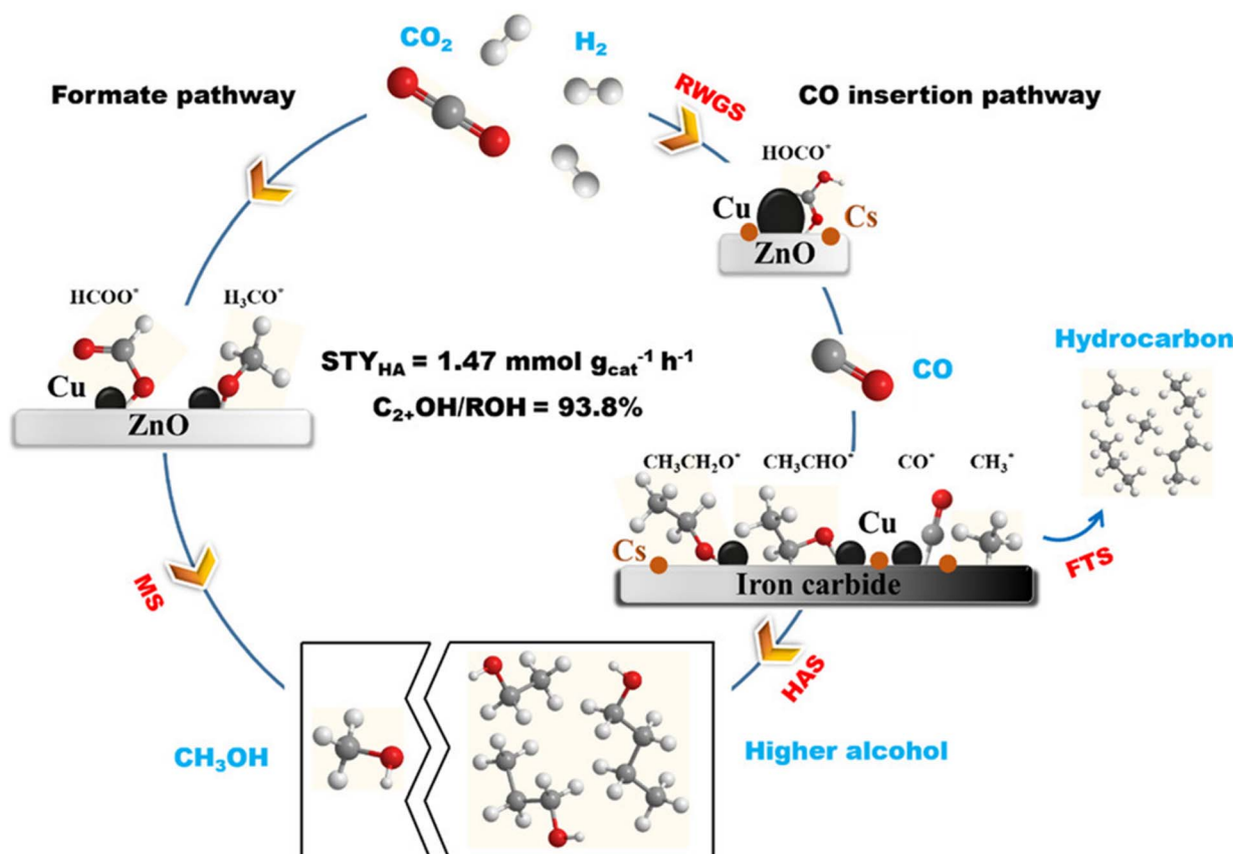


Fig. 10 Proposed reaction mechanism of CO₂ hydrogenation for higher alcohols through C(H)O* insertion and CO insertion into hydrocarbons over Cs–CuFeZn catalysts. Reprinted with permission from ref. 37. Copyright 2020 American Chemical Society.



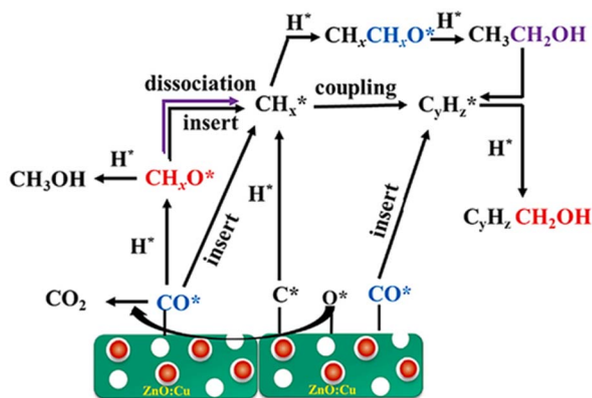


Fig. 11 Electronically modified Fermi level facilitates the formation of alkyl species to promote CO insertion with $\text{CH}_x\text{-C}(\text{H}_x)\text{O}$ coupling for higher alcohols. Reprinted with permission from ref. 17. Copyright 2023 Elsevier.

HCOO^* intermediates following the formate pathway. A proper proportion of the Cs promoter reduces the rate of methanol formation, which is favorable to HAS. Furthermore, at high temperatures, particularly above 300 °C, thermodynamics limits the reaction equilibrium and causes methanol to convert back to CO_2 and H_2 , while the formation of CO is considerably accelerated. However, at high temperatures, the formation of higher alcohols is more favorable because Cu–ZnO produces CO through the RWGS reaction. At the same time, copper–iron carbide facilitates the insertion of $\text{C}(\text{H})\text{O}^*$ species to surface hydrocarbon moieties to produce C_{2+}OH at a similar rate. Furthermore, the Cs promoter is essential for driving the CO insertion reaction during the CO_2 -to-HA process by regulating

the hydrogenation reaction over the CuFeZn catalyst. Since it still does not reach its thermodynamic equilibrium, the rate of C_{2+}OH formation increases as temperature increases. Likewise, a large $\text{C}_{2+}\text{OH}/\text{ROH}$ proportion is produced at high temperatures because of the comparatively slow rate of methanol synthesis in conjunction with the increased rate of C_{2+}OH formation.

Li *et al.*¹⁷ proposed the reaction pathway for HAS from CO_2 hydrogenation over an electronically modified ZnO catalyst, as illustrated in Fig. 11. According to the analysis of the characterization and the activity of an electronically modified ZnO catalyst, the synergistic effect between ZnO and Cu drives the catalytic activity of CO_2 hydrogenation, which helps to propose the reaction mechanism for HAS. ZnO plays a critical role in facilitating the activation of CO and H_2 and promoting the dissociation of the C–O bond. Moreover, the oxygen deficiency species (Zn^{x+} , $0 < x < 2$) promotes carbon chain growth in ZnO. The non-dissociative hydrogenation of CO to generate methoxy species is facilitated by Cu embedded on the ZnO surface. Furthermore, Cu controls the regulation of the electronically modified Fermi level within the ZnO lattice and thus accelerates the electron transfer, facilitating the formation of surface alkyl species (CH_x) by direct C–O dissociation or H-assisted C–O dissociation. As a result, the mechanism of CO insertion with $\text{CH}_x\text{-C}(\text{H}_x)\text{O}$ coupling produces higher alcohols (C_{2+}).

Ye *et al.*⁵³ proposed the reaction pathway for ethanol production from CO_2 hydrogenation over the $\beta\text{-Mo}_2\text{C}$ nanowires with a single Rh atom and an alkali metal K ($\text{K}_{0.2}\text{Rh}_{0.2}/\beta\text{-Mo}_2\text{C}$), as illustrated in Fig. 12. An unmodified $\beta\text{-Mo}_2\text{C}$ catalyst could not convert CO_2 into ethanol (or limit the conversion to only methanol). Rh loading on the $\beta\text{-Mo}_2\text{C}$ catalyst promoted the dissociative adsorption of CO_2 , and the CO^* was adsorbed on

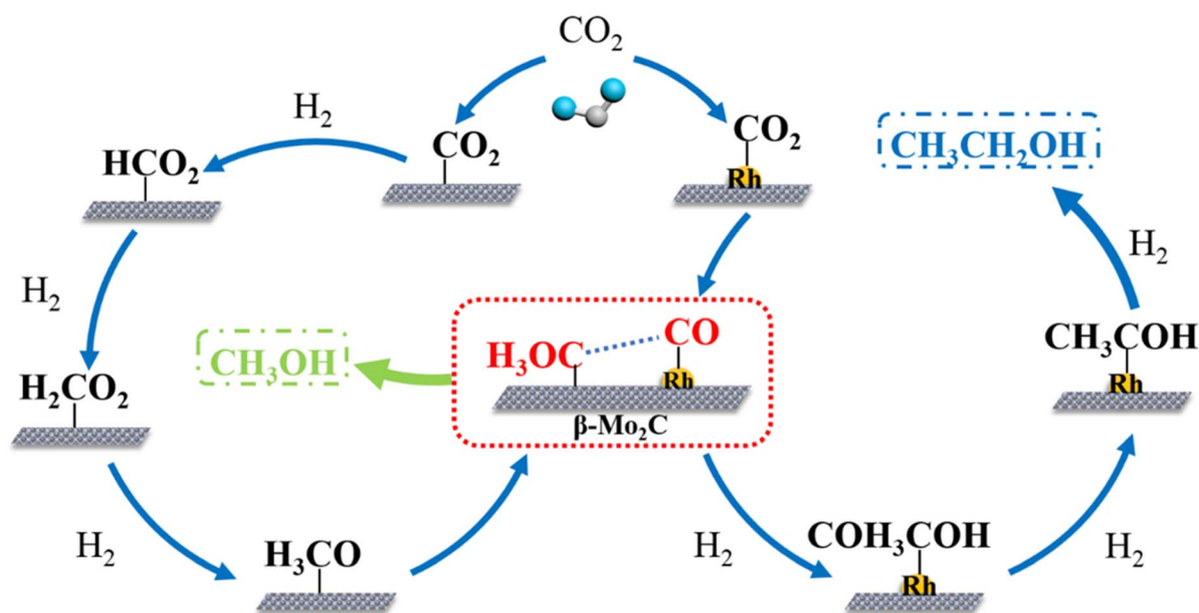


Fig. 12 Proposed reaction mechanism of CO_2 hydrogenation to ethanol on the $\beta\text{-Mo}_2\text{C}$ nanowires with a single Rh atom for promoting the C–C coupling mechanism and an introduction of an alkali metal K to regulate the balanced performance of the two active centers for improving the selectivity of ethanol. Reprinted with permission from ref. 53. Copyright 2022 Elsevier.



the monoatomic Rh^{δ+}. Moreover, hydrogen was more likely to be adsorbed and activated on Rh, which helped to prevent the production of methanol and methane. Additionally, Mo^{δ+}-CO, Mo²⁺-CO, and Mo⁴⁺-CO exhibited IR peaks at 2094, 2108, and 2129 cm⁻¹, respectively. The peaks at 1081, 1053, and 1033 cm⁻¹ were assigned to strongly adsorbed carbonate species (CO₃^{δ-}) formed on the Rh_{1.0}/βMo₂C. Thus, the bifunctional catalyst of monoatomic Rh on βMo₂C was tailored for promoting the C-C coupling. However, achieving high ethanol selectivity by merely adjusting the Rh content was challenging, as increased Rh led to changes in its dispersion and enhanced oxidation of the carrier, resulting in dominant methanol production under mild conditions. Effective C-C coupling relied on SAC bifunctional active centers, where optimized synergy between the two catalytic pathways was essential for efficient carbon chain growth. To further improve the selectivity of ethanol, introducing K assisted in the adsorption and activation of CO₂ on the βMo₂C surface by generating more carbonate species. Furthermore, adding an appropriate amount of K assisted in regulating the carrier properties by controlling the H₂ activation, which could effectively reduce the formation of methanol, yielding a higher selectivity of ethanol. In addition, the nature of single atom Rh was retained to promote the C-C coupling, resulting in ethanol formation.

Conclusions and future perspectives

CO₂ hydrogenation to higher alcohols is a novel strategy to mitigate CO₂ emissions. Efforts have been made to improve the CO₂ conversion and resolve the unsatisfactory selectivity towards higher alcohols. The stability of the catalyst is also a key issue in CO₂ hydrogenation due to catalyst deactivation. The construction of catalysts with various combinations of metals and metal oxides has garnered attention and emerged as a rising trend in enhancing HAS. The introduction of second active metal sites has been proven to hydrogenate the reaction to higher alcohols instead of C₁ product formation, such as methane and methanol. However, designing an ideal catalyst is not easy to achieve. It requires precise control of interplay between active metal sites, supports, and promoters, which could assist in enhanced dispersion of the active metal sites, modify electronic properties, assist in the adsorption and activation of CO₂ on the surface of catalysts, control the activation of H₂, and promote the carbon chain growth, which could suppress the formation of methane and methanol and thus enhancing the activity and selectivity towards higher alcohols. These factors induce a synergistic interaction among the active metal, the support, and the promoter to facilitate a higher conversion of CO₂ with enhanced selectivity for higher alcohols, as discussed in detail in the case-by-case studies presented in this review. Our understanding remains limited despite significant efforts over the past few years to study catalyst design for product selectivity control and mechanistic investigation in HAS. Unlike the well-established and commercially industrialized C₁ production, such as methanol synthesis, the complexity of these reaction pathways and the challenges in catalyst design (requiring a balance among product selectivity, catalyst

stability, and catalytic activity) pose substantial hurdles. Therefore, in-depth investigations of the nature of active sites and reaction pathways through combining advanced characterization (*e.g.*, *in situ* XPS, *in situ* DRIFTS, and CO-TPSR-MS) with computational catalysts (*e.g.*, DFT calculations and machine learning) are strongly recommended to develop highly efficient and cost-effective catalysts. After all, future trends in catalyst design can focus on tailoring the interaction between the active metal sites and the support through the formation of strong metal support interaction (SMSI) to optimize the catalytic activity of catalysts, adjusting the appropriate amount of promoter to enhance the carbon chain growth, prevent the formation of C₁ products, and boost the selectivity towards higher alcohols. Additionally, promoting the confinement effect of supports to reduce particle size and understanding the reaction mechanism to predict the catalytic reaction pathway are essential for achieving a higher conversion of CO₂ and greater selectivity for higher alcohols.

Data availability

The data supporting the findings in this study are sourced from existing literature. Citations are given throughout the paper. Readers may contact the corresponding author, A. C. K. Y., to inquire about this review paper.

Author contributions

A. T.: methodology, investigation, data curation, formal analysis, writing – original draft, writing – review & editing; M. I.: methodology, investigation, data curation, formal analysis, writing – original draft, writing – review & editing; M. P.: formal analysis, writing – original draft, writing – review & editing, funding acquisition; A. C. K. Y.: methodology, investigation, data curation, formal analysis, writing – original draft, writing – review & editing, project administration.

Conflicts of interest

There are no conflicts to declare.

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References

- 1 J. Zhong, X. Yang, Z. Wu, B. Liang, Y. Huang and T. Zhang, *Chem. Soc. Rev.*, 2020, **49**, 1385–1413.
- 2 J. Rogelj, D. Huppmann, V. Krey, K. Riahi, L. Clarke, M. Gidden, Z. Nicholls and M. Meinshausen, *Nature*, 2019, **573**, 357–363.
- 3 Y. Gao, X. Gao and X. Zhang, *Engineering*, 2017, **3**, 272–278.
- 4 G. Yergaziyeva, Z. Kuspanov, M. Mambetova, N. Khudaibergenov, N. Makayeva and C. Daulbayev, *J. CO₂ Util.*, 2024, **80**, 102682.



- 5 R.-P. Ye, J. Ding, W. Gong, M. D. Argyle, Q. Zhong, Y. Wang, C. K. Russell, Z. Xu, A. G. Russell, Q. Li, M. Fan and Y.-G. Yao, *Nat. Commun.*, 2019, **10**, 5698.
- 6 Y. Jiang, K. Wang, Y. Wang, Z. Liu, X. Gao, J. Zhang, Q. Ma, S. Fan, T.-S. Zhao and M. Yao, *J. CO₂ Util.*, 2023, **67**, 102321.
- 7 S. Aslam, S. Rani, K. Lal, M. Fatima, T. Hardwick, B. Shirinfar and N. Ahmed, *Green Chem.*, 2023, **25**, 9543–9573.
- 8 I. Dincer, *Int. J. Hydrogen Energy*, 2012, **37**, 1954–1971.
- 9 A. Álvarez, A. Bansode, A. Urakawa, A. V. Bavykina, T. A. Wezendonk, M. Makkee, J. Gascon and F. Kapteijn, *Chem. Rev.*, 2017, **117**, 9804–9838.
- 10 P. Gao, L. Zhang, S. Li, Z. Zhou and Y. Sun, *ACS Cent. Sci.*, 2020, **6**, 1657–1670.
- 11 F. Dalena, A. Senatore, M. Basile, S. Knani, A. Basile and A. Iulianelli, *Membranes*, 2018, **8**, 98.
- 12 M. A. Ratcliff, J. Luecke, A. Williams, E. Christensen, J. Yanowitz, A. Reek and R. L. McCormick, *Environ. Sci. Technol.*, 2013, **47**, 13865–13872.
- 13 V. F. Andersen, J. E. Anderson, T. J. Wallington, S. A. Mueller and O. J. Nielsen, *Energy Fuels*, 2010, **24**, 3647–3654.
- 14 D. Xu, Y. Wang, M. Ding, X. Hong, G. Liu and S. C. E. Tsang, *Chem*, 2021, **7**, 849–881.
- 15 J. M. Spero, B. DeVito and L. Theodore, *Regulatory Chemicals Handbook*, CRC Press, 1st edn, 2000, DOI: [10.1201/9781482270389](https://doi.org/10.1201/9781482270389).
- 16 C. Yang, B. Wang, Y. Wen, M. Fan, Y. Jia, S. Zhou and W. Huang, *Fuel*, 2022, **327**, 125055.
- 17 F. Li, P. Jia, Q. Zhang, Y. Liu, V. A. Vinokurov and W. Huang, *Fuel Process. Technol.*, 2023, **241**, 107600.
- 18 G. Cui, Y. Lou, M. Zhou, Y. Li, G. Jiang and C. Xu, *Catalysts*, 2024, **14**, 232.
- 19 A. I. Latsiou, N. D. Charisiou, Z. Frontistis, A. Bansode and M. A. Goula, *Catal. Today*, 2023, **420**, 114179.
- 20 D. Xu, M. Ding, X. Hong and G. Liu, *ACS Catal.*, 2020, **10**, 14516–14526.
- 21 B. An, Z. Li, Y. Song, J. Zhang, L. Zeng, C. Wang and W. Lin, *Nat. Catal.*, 2019, **2**, 709–717.
- 22 J. Nebel, S. Schmidt, Q. Pan, K. Lotz, S. Kaluza and M. Muhler, *Chin. J. Catal.*, 2019, **40**, 1731–1740.
- 23 R. Yao, J. Wei, Q. Ge, J. Xu, Y. Han, Q. Ma, H. Xu and J. Sun, *Appl. Catal., B*, 2021, **298**, 120556.
- 24 R. Yao, B. Wu, Y. Yu, N. Liu, Q. Niu, C. Li, J. Wei and Q. Ge, *Appl. Catal., B*, 2024, **355**, 124159.
- 25 Y. Wang, K. Wang, B. Zhang, X. Peng, X. Gao, G. Yang, H. Hu, M. Wu and N. Tsubaki, *ACS Catal.*, 2021, **11**, 11742–11753.
- 26 Y. Zhou, Y. Wang, H. Lu, T. Liu, X. Hong and G. Liu, *ACS Sustain. Chem. Eng.*, 2024, **12**, 3322–3330.
- 27 L. Wang, L. Wang, J. Zhang, X. Liu, H. Wang, W. Zhang, Q. Yang, J. Ma, X. Dong, S. J. Yoo, J. G. Kim, X. Meng and F. S. Xiao, *Angew Chem. Int. Ed. Engl.*, 2018, **57**, 6104–6108.
- 28 J. Graciani, D. C. Grinter, P. J. Ramírez, R. M. Palomino, F. Xu, I. Waluyo, D. Stacchiola, J. Fdez Sanz, S. D. Senanayake and J. A. Rodriguez, *ACS Catal.*, 2022, **12**, 15097–15109.
- 29 G. Wang, R. Luo, C. Yang, J. Song, C. Xiong, H. Tian, Z.-J. Zhao, R. Mu and J. Gong, *Sci. China: Chem.*, 2019, **62**, 1710–1719.
- 30 B. Ouyang, S. Xiong, Y. Zhang, B. Liu and J. Li, *Appl. Catal., A*, 2017, **543**, 189–195.
- 31 D. Wang, Q. Bi, G. Yin, W. Zhao, F. Huang, X. Xie and M. Jiang, *Chem. Commun.*, 2016, **52**, 14226–14229.
- 32 M. Kishida, K. Yamada, H. Nagata and K. Wakabayashi, *Chem. Lett.*, 2006, **23**, 555–556.
- 33 B. Kang, S. Qi, X. Wang, F. Bai, N. Wang, R. Hu, Y. Zhang and H. Su, *Catal. Commun.*, 2020, **137**, 105945.
- 34 S. Liu, H. Zhou, Q. Song and Z. Ma, *J. Taiwan Inst. Chem. Eng.*, 2017, **76**, 18–26.
- 35 L. Ding, T. Shi, J. Gu, Y. Cui, Z. Zhang, C. Yang, T. Chen, M. Lin, P. Wang, N. Xue, L. Peng, X. Guo, Y. Zhu, Z. Chen and W. Ding, *Chem*, 2020, **6**, 2673–2689.
- 36 M. Irshad, H.-J. Chun, M. K. Khan, H. Jo, S. K. Kim and J. Kim, *Appl. Catal., B*, 2024, **340**, 123201.
- 37 D. Xu, M. Ding, X. Hong, G. Liu and S. C. E. Tsang, *ACS Catal.*, 2020, **10**, 5250–5260.
- 38 D. Xu, H. Yang, X. Hong, G. Liu and S. C. Edman Tsang, *ACS Catal.*, 2021, **11**, 8978–8984.
- 39 Q. Zhang, S. Wang, R. Geng, P. Wang, M. Dong, J. Wang and W. Fan, *Appl. Catal., B*, 2023, **337**, 123013.
- 40 S. Zhang, Z. Wu, X. Liu, Z. Shao, L. Xia, L. Zhong, H. Wang and Y. Sun, *Appl. Catal., B*, 2021, **293**, 120207.
- 41 K. An, S. Zhang, J. Wang, Q. Liu, Z. Zhang and Y. Liu, *J. Energy Chem.*, 2021, **56**, 486–495.
- 42 X. Ding, J. Fu, Y. Lyu, L. Ma, Y. Xu and X. Liu, *Chem. Eng. J.*, 2024, **494**, 152923.
- 43 T. Wittoon, T. Numpilai, S. Nijpanich, N. Chanlek, P. Kidkhunthod, C. K. Cheng, K. H. Ng, D.-V. N. Vo, S. Ittisanronnachai, C. Wattanakit, M. Chareonpanich and J. Limtrakul, *Chem. Eng. J.*, 2022, **431**, 133211.
- 44 X. Xi, F. Zeng, H. Zhang, X. Wu, J. Ren, T. Bisswanger, C. Stampfer, J. P. Hofmann, R. Palkovits and H. J. Heeres, *ACS Sustain. Chem. Eng.*, 2021, **9**, 6235–6249.
- 45 H. Yang, Z. Wei, J. Zhang, Y. Dang, S. Li, X. Bu, Z. Zhou, C. Gong, H. Wang, J. Li, Y. Liu, Y. Yang, T. Xiao, C. Liu, Y. Sun and P. Gao, *Chem*, 2024, **10**, 2245–2265.
- 46 X. Ye, C. Yang, X. Pan, J. Ma, Y. Zhang, Y. Ren, X. Liu, L. Li and Y. Huang, *J. Am. Chem. Soc.*, 2020, **142**, 19001–19005.
- 47 Y. Lou, F. jiang, W. Zhu, L. Wang, T. Yao, S. Wang, B. Yang, B. Yang, Y. Zhu and X. Liu, *Appl. Catal., B*, 2021, **291**, 120122.
- 48 S. Bai, Q. Shao, P. Wang, Q. Dai, X. Wang and X. Huang, *J. Am. Chem. Soc.*, 2017, **139**, 6827–6830.
- 49 Y. Wang, Y. Zhou, X. Zhang, M. Wang, T. Liu, J. Wei, G. Zhang, X. Hong and G. Liu, *Appl. Catal., B*, 2024, **345**, 123691.
- 50 F. Zhang, W. Zhou, X. Xiong, Y. Wang, K. Cheng, J. Kang, Q. Zhang and Y. Wang, *J. Phys. Chem. C*, 2021, **125**, 24429–24439.
- 51 S. Ji, F. Hong, D. Mao, Q. Guo and J. Yu, *Chem. Eng. J.*, 2024, **495**, 153633.
- 52 C.-C. Tran and S. Kaliaguine, *Chem. Eng. J.*, 2024, **496**, 153636.
- 53 X. Ye, J. Ma, W. Yu, X. Pan, C. Yang, C. Wang, Q. Liu and Y. Huang, *J. Energy Chem.*, 2022, **67**, 184–192.
- 54 H. Lei, R. Nie, G. Wu and Z. Hou, *Fuel*, 2015, **154**, 161–166.
- 55 H. Lei, Z. Hou and J. Xie, *Fuel*, 2016, **164**, 191–198.



- 56 W. Wang, S. Wang, X. Ma and J. Gong, *Chem. Soc. Rev.*, 2011, **40**, 3703.
- 57 Y. Suo, Y. Yao, Y. Zhang, S. Xing and Z.-Y. Yuan, *J. Ind. Eng. Chem.*, 2022, **115**, 92–119.
- 58 J. Graciani, K. Mudiyansele, F. Xu, A. E. Baber, J. Evans, S. D. Senanayake, D. J. Stacchiola, P. Liu, J. Hrbek, J. F. Sanz and J. A. Rodriguez, *Science*, 2014, **345**, 546–550.
- 59 C. Yang, S. Liu, Y. Wang, J. Song, G. Wang, S. Wang, Z.-J. Zhao, R. Mu and J. Gong, *Angew. Chem., Int. Ed.*, 2019, **58**, 11242–11247.
- 60 M. Zhang, R. Yao, H. Jiang, G. Li and Y. Chen, *Appl. Surf. Sci.*, 2017, **412**, 342–349.
- 61 S. Liu, C. Yang, S. Zha, D. Sharapa, F. Studt, Z.-J. Zhao and J. Gong, *Angew. Chem., Int. Ed.*, 2022, **61**, e202109027.
- 62 G. Prieto, *ChemSusChem*, 2017, **10**, 1056–1070.
- 63 N. A. Sholeha, H. Holilah, H. Bahruji, A. Ayub, N. Widiastuti, R. Ediati, A. A. Jalil, M. Ulfa, N. Masruchin, R. E. Nugraha and D. Prasetyoko, *S. Afr. J. Chem. Eng.*, 2023, **44**, 14–30.
- 64 M. Wang, P. Wang, G. Zhang, Z. Cheng, M. Zhang, Y. Liu, R. Li, J. Zhu, J. Wang, K. Bian, Y. Liu, F. Ding, T. P. Senftle, X. Nie, Q. Fu, C. Song and X. Guo, *Sci. Adv.*, 2023, **9**, eadg0167.
- 65 H. Kusama, K. Okabe, K. Sayama and H. Arakawa, *Catal. Today*, 1996, **28**, 261–266.
- 66 K. Kitamura Bando, K. Soga, K. Kunimori and H. Arakawa, *Appl. Catal., A*, 1998, **175**, 67–81.
- 67 X. Li, M. Song, Y. Zhou, P. Zhou, D. Xu, T. Liu and X. Hong, *ChemCatChem*, 2024, **16**, e202301577.
- 68 H. Xin, L. Lin, R. Li, D. Li, T. Song, R. Mu, Q. Fu and X. Bao, *J. Am. Chem. Soc.*, 2022, **144**, 4874–4882.
- 69 W. Zhang, H. Lin, Y. Wei, X. Zhou, Y. An, Y. Dai, Q. Niu, T. Lin and L. Zhong, *ACS Catal.*, 2024, **14**, 2409–2417.
- 70 M. Ren, Y. Zhang, X. Wang and H. Qiu, *Catalysts*, 2022, **12**, 403.
- 71 S. Zhang, X. Liu, Z. Shao, H. Wang and Y. Sun, *J. Catal.*, 2020, **382**, 86–96.
- 72 J. Wang, R. Li, G. Zhang, C. Dong, Y. Fan, S. Yang, M. Chen, X. Guo, R. Mu, Y. Ning, M. Li, Q. Fu and X. Bao, *J. Am. Chem. Soc.*, 2024, **146**, 5523–5531.
- 73 D. Iltsiou, J. Mielby and S. Kegnaes, *Chempluschem*, 2024, **89**, e202300313.
- 74 B. Liu, B. Ouyang, Y. Zhang, K. Lv, Q. Li, Y. Ding and J. Li, *J. Catal.*, 2018, **366**, 91–97.
- 75 S. Zhang, K. An, J. Xin, Y. Jiang, M. Niu, P. Song, C. Wu, H. Wang and Y. Liu, *Chem. Eng. J.*, 2024, **486**, 150334.
- 76 C. Zhang, L. Wang, U. J. Etim, Y. Song, O. M. Gazit and Z. Zhong, *J. Catal.*, 2022, **413**, 284–296.
- 77 T. Liu, D. Xu, M. Song, X. Hong and G. Liu, *ACS Catal.*, 2023, **13**, 4667–4674.
- 78 S. Liu, Y. He, W. Fu, J. Chen, J. Ren, L. Liao, R. Sun, Z. Tang, C. Mebrahtu and F. Zeng, *J. CO2 Util.*, 2023, **67**, 102322.

