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Recent advances in CO₂ hydrogenation to methane using single-atom catalysts

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Single-atom catalysts (SACs) with isolated metal atoms dispersed on a solid support have been a hot topic in the last decade in heterogeneous catalysis. In a comprehensive attempt to minimize CO₂ emissions with a persisting effect on global warming, the catalytic conversion of CO₂ to methane can facilitate the convenient storage of excess energy as a green natural gas. Herein, we narrow the scope of this review by focusing on recent reports on thermocatalytic CO₂ methanation using SACs with a special focus on the design and optimization strategies and existing challenges to overcome to get high selectivity and activity. It is highlighted that for higher selectivity towards methane over the CO product in CO₂ hydrogenation, H₂ dissociation and strong CO binding as an intermediate over the catalyst surface are the key steps. These insights are vital for determining the design principles of upcoming SACs to make them more efficient and selective in CO₂ methanation.

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Sustainability spotlight

Single-atom catalysts (SACs) are among the most important tools of the new era for commercializing CO₂ emissions. Among CO₂RR products, methane (CH₄), being a renewable fuel hydrocarbon, has attracted significant attention, and its storage, transportation, and applications have been well researched. The invention of SACs on the whole greatly alleviates the obstacle of scaling-up processes that previously used precious metals as catalysts that are highly selective and effective toward CO₂ methanation. In this review, the issue of product selectivity in CO₂ hydrogenation, with a focus on mentioning the synthesis strategies of SACs and their utilization for CO₂ methanation reactions, is highlighted. We strongly believe that this review will enlighten ongoing and future catalyst research, offering fundamental guidance to design multifarious SACs for CO₂ methanation.

1. Introduction

The global demand for modern material designs with redefined catalytic efficiency is driving the circular economy to new heights. As the need for carbon neutrality intensifies, newly invented catalysts for the conversion of CO₂ by CO₂ reduction reactions (CO₂RR) or CO₂ hydrogenation to value-added chemicals must be able to surpass the bottlenecks of the existing catalytic chemistry.^{1,2} Based on the actual requirements, diverse subcategories and plentiful designs of heterogeneous catalysts have been developed and applied by global industries. The most explored classes include metal oxides,^{3–8} metal sulfides,^{9–11} metal–organic frameworks (MOFs),^{12–17} covalent organic frameworks (COFs),^{18,19} carbonaceous materials²⁰ (such as carbon nanofibers,^{21–24} graphene,^{25,26} and carbon nanotubes (CNTs)^{27–29}), and conjugated microporous polymers (CMPs).³⁰ In this context, single-atom catalysts (SACs) are one of the most promising classes of catalysts, and have shown good

potential due to offering high catalytic activity and a high exposure to active sites together with good recyclability and reusability.

Recently, SACs have been receiving much attention from the scientific community owing to their extraordinary properties of both homogeneous and heterogeneous catalysts (Fig. 1). SACs can solve the common issues of homogeneous catalysts, in term of their recyclability and high cost and difficult synthetic procedure, without compromising the product selectivity.^{31,32} Since SACs are considered one of the most important tools of the new era for commercializing CO₂ emissions, considerable efforts have been made in the last decades to achieve the selective hydrogenation of CO₂ for ensuring the desired products.^{33–37} Among CO₂RR products, methane (CH₄), being a renewable fuel hydrocarbon, has attracted much attention, and its storage, transportation, and applications have been well researched. To date, it has been reported that conventional catalysts can achieve a maximum 50% selectivity, but with an unsatisfactory practical current density (>100 mA cm⁻²).³³ Herein, we highlight the issue of product selectivity in CO₂ hydrogenation by considering the synthesis strategies of SACs and their utilization for CO₂ methanation reactions.

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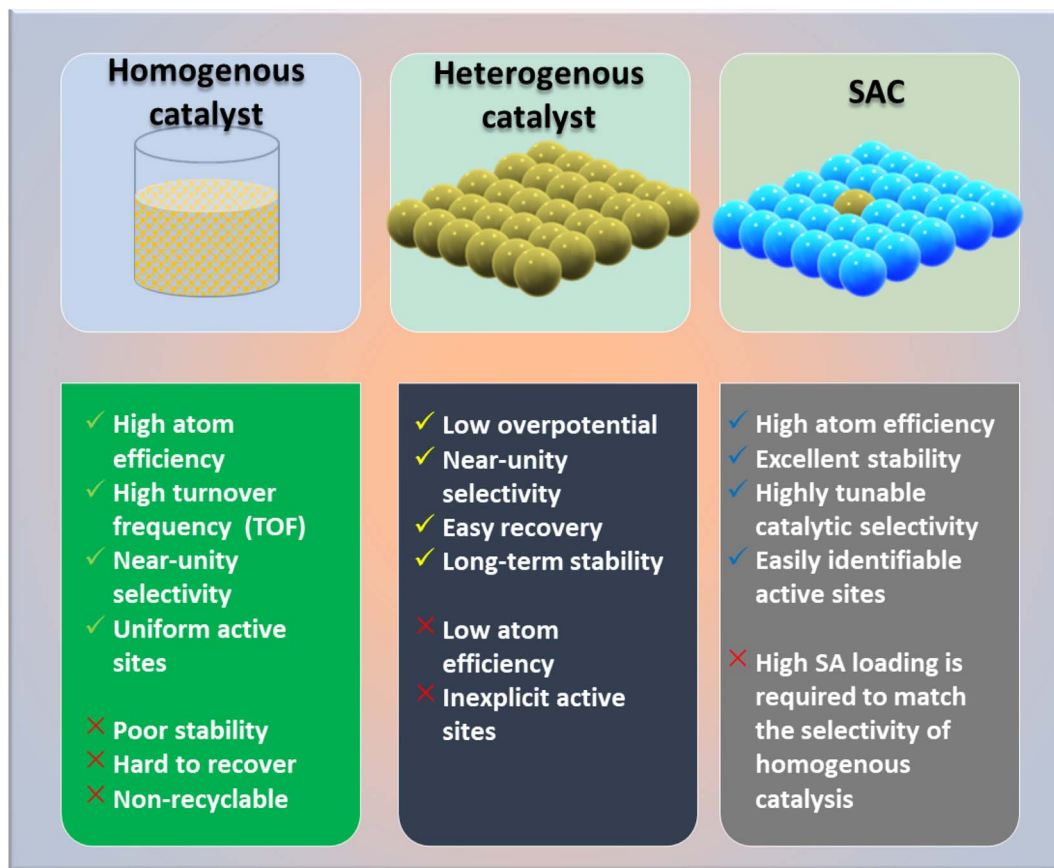


Fig. 1 Comparison of the properties of SACs with those of homogeneous and heterogeneous catalysts.

1.1. Scope of the review

Owing to the meteoric rise in publications in SACs over the last few years, several attempts were previously made to review SACs, including the fundamentals,^{38,39} various advancements made in designing SACs,³⁷ and theoretical insights⁴⁰ into the electronic structure and mechanistic aspects of catalysis in various organic reactions.⁴¹ Many reviews have discussed the scope of SACs for CO₂ hydrogenation transformation alone.^{36,42–50} Quite a large number of progress reports have also been made on SACs for the electrochemical reduction of CO₂.^{36,42–47} Herein, we narrow the scope of this review by focusing on CO₂RR methanation using SACs in light of relevant reports. For the sake of understanding, we provide a brief overview of the general aspects of SACs, including the theory, their structures, and common synthetic strategies. Afterward, we discuss how SACs have been employed for CO₂ hydrogenation into useful products, further narrowing the discussion down to the methanation of CO₂ in particular. Additionally, an earnest attempt is made to give the reader a comprehensive understanding of the mechanistic aspects of SAC methanation. It is remarkable that around one-third of reports on SACs are based on theoretical conceptualization,⁴⁰ while a majority of the experimental reports endorsed their results based on theoretical grounds. We strongly believe that this review will enlighten ongoing and future catalyst research, offering fundamental guidance to aid the design of multifarious SACs for CO₂ methanation.

1.2. SACs: evolution, structure, and theory

Numerous studies and observations have shown that the efficiency and selectivity of a heterogeneous catalyst are directly related to its size.^{51–56} It is evident that as a catalyst's size is decreased from the micron to cluster to atomic size, more catalytic access points are created or more active sites can be accessed, which leads to an increase in catalytic efficiency.^{57,58} It is noteworthy that even though downsizing catalysts into (sub)nanoclusters is highly desirable for catalysis, stabilizing such systems under realistic reaction conditions still requires significant effort, despite the advancements accomplished in nanotechnology. For example, metal nanoclusters characteristically display instability, especially under thermocatalytic conditions, which prompts sintering them into larger particles (such as nanoparticles, and nano aggregates).^{59–61} However, over time, it has been established that these clusters could better withstand extreme conditions by encapsulating them into molecular frameworks or cage-like structures, like zeolites, COFs, and MOFs.⁵⁹ Nevertheless, reducing a catalytic active element to the atomic level (equivalent to a homogenous catalyst) can lead to improvements in catalytic efficiency, but these would still be highly unstable under real-time reaction conditions. This is because free atoms or active elements with unsaturated coordination sites are too reactive.⁶² Moreover, it has been revealed that the existence of an isolated atom can restrain such a risk when it is bound to a suitable support in



a crystal lattice. A heterogeneous catalyst so formulated is called a single-atom catalyst (SAC) and can overcome many of the drawbacks of a typical heterogeneous catalyst.

The term 'single-atom catalyst' was first coined by Qiao *et al.*⁶² in 2011. SACs can be defined as an ensemble of any catalytically active atom spatially secluded from its own species by directly bonding with a solid carrier host of another entity. The terminology SAC should not be confused with the isolated active elements immobilized with the linker functionalities of carriers.⁶³ It is worth mentioning that, even before introducing the terminology SAC, many groups had already reported the exceptional catalytic features of SAC systems but variously termed these as 'isolated atom', 'atomically dispersed', or 'single site catalysts', where the single metal atoms were isolated on a metal oxide support.^{64,65} Seemingly, SACs stand as a bridge between both types of catalysts. A heterogeneous catalyst has a bad reputation for its less viable active sites for molecular transformation. Unlike a homogenous catalyst, a heterogeneous catalyst, however, offers better thermal stability and easy recovery. SACs provide a solid insoluble platform that is viable for easy recovery with enough chemical stability but with possibly 100% exposed active sites. The evolution of SACs right after heterogeneous catalysts is illustrated in Fig. 2.^{62,66–75}

Typically, the design of SACs has three main components: (i) active single atom (SA) entity, (ii) SAC support, and (iii) anchoring sites and anchoring strategies, as shown in Fig. 3. Being highly selective and effective for CO₂ methanation, the active SA entity of SACs is the part responsible for the catalytic action. Ru, Rh, and Pd are the most explored noble metals and are known for the selective methanation of CO₂.^{76,77} A prudent and maximal utilization of these SAs in SACs would help economize CO₂ methanation to a significant extent by reducing the overall consumption of precious metals. Besides, providing a proper coordination environment and electronic structure of a SA could, on the other hand, also address the limited activity and deactivation issue of low-cost metals, like Ni, toward the methanation of CO₂. The SAC support, as the second major

component of SACs, helps to stabilize the SA entity. The stability of the SA in SACs largely depends on how the SA is bonded to the support. Typically, there are 3 classes of support: metal oxides, carbon materials, and porous structures (zeolites, MOF, COF, *etc.*). The porous structures help to significantly increase SA loading onto the support matrix.⁷⁸

The third designating entity in SACs is their anchoring strategy, which plays a decisive role in the SAC accomplishing a particular reaction out of a certain SA-support combination. The anchoring of the SA onto the support could be achieved using dopants, defects, or confinements. The heteroatom dopants in SACs can enhance the catalytic activity by creating surface defects to stabilize the structure.^{79–81} In such cases, single atom stabilization usually occurs through covalent/ionic interactions with the support surface.^{33,50} Doping can be done by introducing non-metals into the lattices or interstitial sites of a host system. Non-metallic elements with lone pairs (such as N, S, or O) doped in a host system can provide anchoring centers for metallic SAs, leading to the stabilization of the SAC.^{82–85} Defect engineering is another strategy to introduce vacancies onto a host material, which otherwise would not be introduced by doping.⁸⁶ Surface step edge defects, unsaturated coordination sites, and cation vacancies are important anchoring centers for single atoms.^{87–89} A high SA loading can also be achieved by confining the active elements spatially in the pores of perforated materials (COFs, zeolites, MOFs) or by linking with coordination sites. For example, 20% atom dispersion was achieved by using melamine sponges with a carbon nitride backbone, which apparently helped to effectively disperse SAs throughout the host system.⁶³ Thus the electronic structure of SACs will be governed by the coordination environment of the SA.⁴⁰ Apparently, the electronic structure of SACs will differ from their bulk counterparts. Since SACs have a regular crystal lattice, simulating equivalent models largely gives insights into the electronic features of the anticipated structure. This is also vital to decide the design principles of SACs to make them selective toward a particular reaction, like the hydrogenation of CO₂.

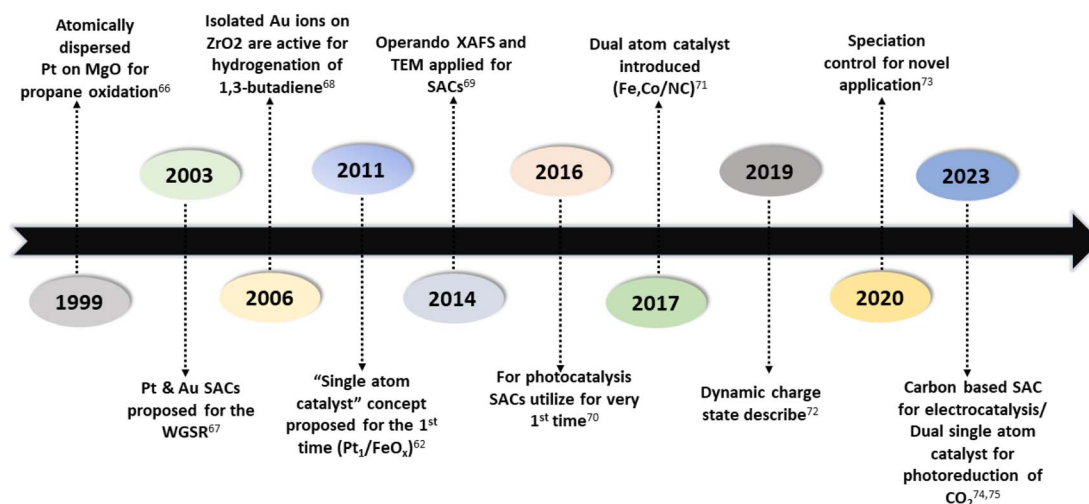


Fig. 2 Evolution of single-atom catalysts over time.



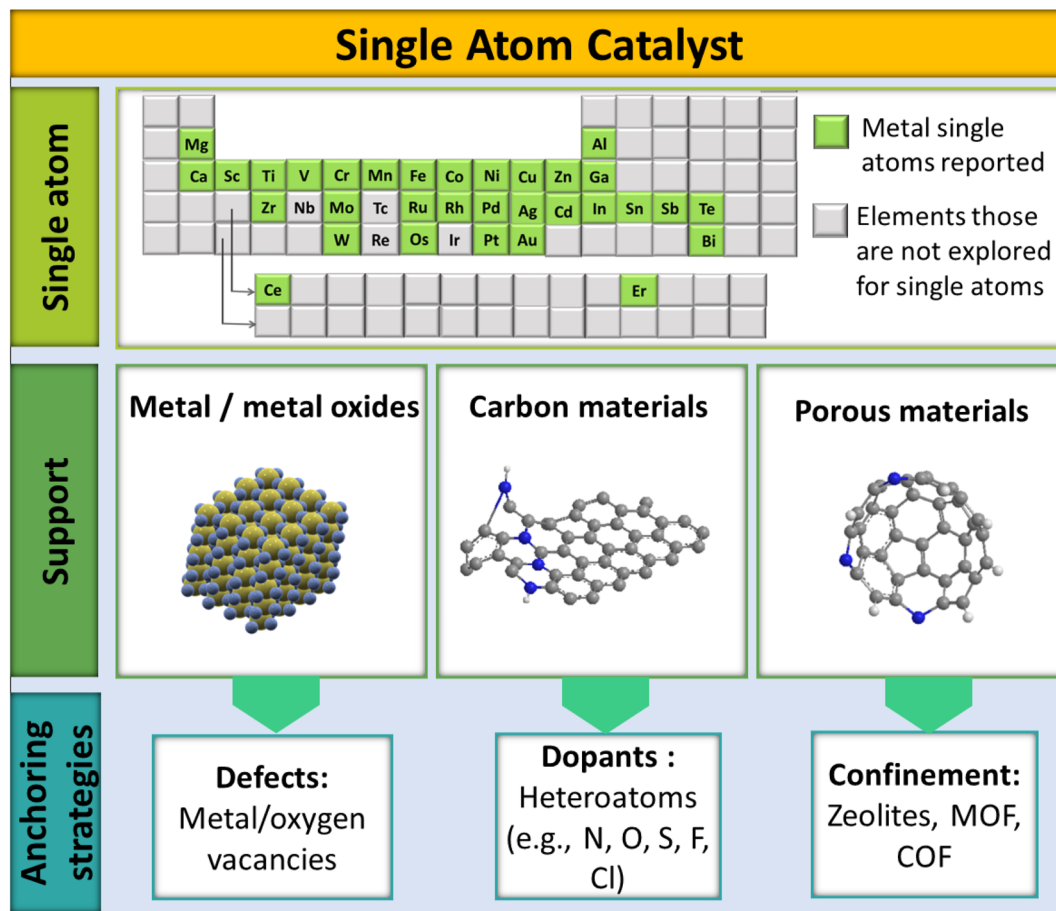


Fig. 3 Illustration of the structural design entities of SACs.

1.3. Synthesis

The bottleneck for acquiring SACs is the controlled growth of a SA on a support phase without compromising the atomic catalytic activity of the active phase. Common strategies that can be adopted to prepare SACs are: (1) lowering the active phase loading, (2) increasing the metal-support interactions, and (3) exploring vacancy defects on the support. Measures like mass-selected soft-landing and atomic layer deposition are helpful in establishing the above scenarios, enabling the precise regulation of the growth of SACs. Nevertheless, scaling up these methods and their economization pose problems. Recently, many wet chemical methods, which are usually simple and easy to implement, have been introduced as alternatives.^{41,62} However, most wet chemical processes rely on typically trial-and-error principles. Altogether, any synthetic routes that already exist or that are being constantly evolved can be classified broadly into bottom-up strategies (direct synthesis) and top-down strategies (post synthesis).

Bottom-up strategies (direct synthesis) are the most common approach to achieve metal SACs. When the minority element precursor is adsorbed, reduced, and/or confined by the vacancy defects or voids present in the host material, single metals can be stabilized in solitary sites by surrounding them with the host material. This can happen during the preparation of the host material. Stabilizing isolated atoms gets the prime focus, which

decides the ultimate thermodynamic stability of SACs. For further in-depth information, we refer to recent reviews dealing with preparation strategies.^{90,91}

In top-down approaches (post synthesis), an isolated single metal of interest is introduced and gets stabilized in the bulk catalyst right after preparation.^{91–94} Many SACs are prepared by incorporating/depositing the active SA in pre-synthesized support having appropriate coordination sites. Common strategies are gas-phase, electrochemical, and wet chemical deposition.¹⁸ More strategies have been constantly evolving over time. The most common methods are discussed below. A concise illustration of the synthetic strategies is presented in Fig. 4. The major merits and demerits of the existing synthesis methods for SACs are listed in Table 1.

(1) Coprecipitation approach: this is a wet chemical method that is known for its straightforwardness. The method needs neither complicated steps nor any special equipment and hence could be executed in any wet chemical laboratory. Here, host precursor are precipitated in parallel with the SA active precursor, resulting in the desired SAC. For example, Lin *et al.* synthesized single-atom Ir species dispersed on a FeO_x support by precipitating from H₂IrCl₆ solution.⁹⁵ In fact, the first identified SAC was prepared *via* a coprecipitation method to isolate Pt atoms dispersed in iron oxide.⁶² Thereafter, researchers devoted efforts to find a reliable root to achieve SACs by



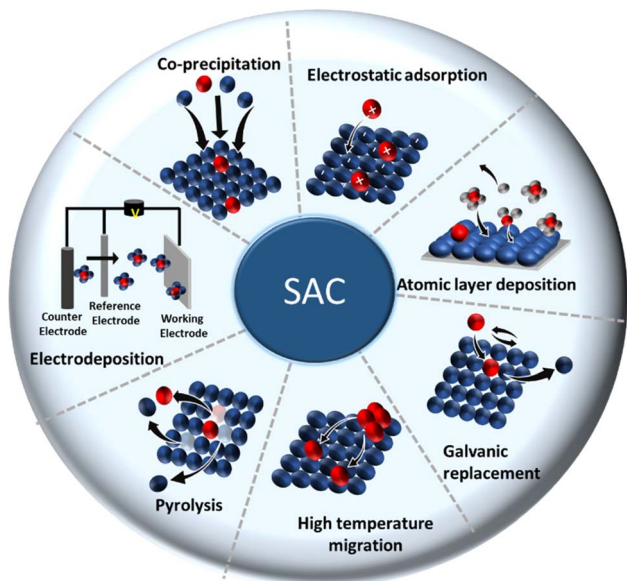


Fig. 4 Pictorial representation of the synthesis methods for SACs.

adopting different coprecipitation strategies suitable for a wide combination of catalytic active elements in a plethora of supports.⁹¹

(2) Electrostatic adsorption: this is a special wet impregnation method, wherein single metal species are adsorbed on the support by a strong electrostatic interaction. Here, a strong metal–support interaction prevents metal–metal interactions, guaranteeing a stable dispersion of the SA phase on the support phase. In some cases, electronic modification of isolated metal atoms by charged ions (cations or anions) helps to stabilize the isolated active species from agglomerating into clusters or nanoparticles. Modification of the precursors and support surface is, therefore, decisive in this method. For example, in SACs, it has been proven that the protection of isolated metal atoms, like Pt and Pd, from aggregation could be accomplished by introducing charged species, like ionic liquids (ILs), on supports, like hydroxyapatite (HAP).⁹⁶ This will not only enhance the kinetic barrier of forming metal–metal bonds but also decrease the thermodynamic feasibility of metal aggregation.

(3) Atomic layer deposition (ALD): in the ALD process, a cyclic self-limiting reaction between a gas-phase precursor and a solid-host surface enables the controlled deposition of single-atom-thick layers one over other.^{97,98} The vapor-phase deposition technique provides a scalable way to achieve SACs with good uniformity and conformity. Due to its high surface chemical selectivity, the possibility of impurities from the residual components is less probable, unlike what happens in the simple coprecipitation method.

(4) Electrodeposition: electrodeposition involves the deposition of an active metal containing an electrolyte onto the solid support. Despite the scalability, economic advantages, and better control over the extent of active element deposition (owing to the control over the applied potential and amount of metal ion precursor in the electrolyte), electrodeposition

possesses a major drawback of a possible inhomogeneous coating onto the host phase.³⁹ Hence, a universal strategy for realizing SACs for a wider class of SA–support combinations is difficult to formulate, but not impossible. Indeed, Zhang *et al.*⁹⁹ demonstrated a universal electrodeposition applicable for the deposition of 4d and 5d metals, like Ru, Rh, Pd, Ag, Pt, and Au, onto a variety of supports, like $\text{Co}(\text{OH})_2$, MnO_2 , MoS_2 , $\text{Co}_{0.8}\text{Fe}_{0.2}\text{Se}_2$ nanosheets, and nitrogen-doped carbon.

(5) Galvanic replacement method: the process for the galvanic replacement reaction (GRR) involves redox reactions, wherein a metal precursor is oxidized, dissolved, and replaced/deposited by the reduction of another metal precursor with a higher reduction potential. Thus, the standard electrode potential of the precursor ions will decide the nature of pairing two components in a particular SAC.⁷⁹ Toshima's group has introduced a crown jewel cluster concept to prepare highly active and scalable colloidal nanocatalyst Au/Pd clusters by adopting this method.^{100,101} Some of the top Pd atoms from Pd nanoclusters (Pd_{147} , considered as a crown) were, thus, galvanically replaced by Au atoms (appeared as a jewel), leaving a high degree of coordinative unsaturation, responsible for high catalytic activity. However, the galvanic replacement in colloidal form often results in dissolution of the non-noble element by depositing the noble metal component. Also, the process is hard to predict, such that the resultant structure of the SAC is governed mainly by the nature of the protective agent and the morphology of the host material.¹⁰² Another drawback of this method is that this process is applicable only to those metals whose electrode reduction potential is higher than that of the host.¹⁰³ Thus, it is speculated that when the reduction potential of the host component is higher than that of the SA component, an alternative way to achieve the SAC or single-atom alloy (SAC) is to downsize the host to promote a reverse replacement reaction.¹⁰³ Besides, it is worth mentioning that galvanic replacement is not only restricted to noble metals but also can be applied for the replacement of transition metals (*e.g.*, Ni), excluding Cu. For example, Peng *et al.*¹⁰⁴ used Ni nanoparticles sized 4.4 nm as a host for the controlled deposition of single Pt atoms using galvanic replacement.

(6) Pyrolysis: this method forms a metal on a C-based host because of the decomposition of the precursor at pyrolysis temperature or by chemical leaching. On the other hand, this strategy involves a sacrificial host template (removed by acid leaching or oxidative calcination). The sacrificial material could be a MOF, COF, zeolite, or those that can hold metals *via* a variety of linkers. A uniformly dispersed SA system with enough thermal stability can be realized by the thermal decomposition of various MOFs. As an alternative, the desired active single metal ion can also be ion exchanged post-synthesis of the MOF, followed by high-temperature pyrolysis.³⁵ Single/bimetallic zeolitic imidazolate frameworks (ZIFs) are the most explored MOFs for preparing SACs for CO_2 reduction. Wang *et al.*³⁴ successfully prepared Co-based SACs by high temperature pyrolysis of Co, Zn-based bimetallic ZIF precursors. Pyrolysis at a temperature above Zn metal's boiling point (*i.e.*, $>907^\circ\text{C}$) facilitated carbonization of the organic linkers, and evaporation of the low boiling Zn, as well as a reduction of



Table 1 Merits and demerits of different types of synthesis strategies reported for SACs

Type of synthesis	Process	Advantages	Disadvantages	Ref.
Coprecipitation	Precipitation of host precursor in parallel with SA precursor	<ul style="list-style-type: none"> • Simple operation • Requires no complicated steps or any special equipment 	<ul style="list-style-type: none"> • Control over single atom loading is less • Heterogeneous active sites • Post-treatments are required to limit the aggregation of isolated SA 	62 and 107
Electrostatic adsorption	Single metal species is adsorbed and stabilized on the support by a strong electrostatic interaction	<ul style="list-style-type: none"> • Simple operation • Scalable 	<ul style="list-style-type: none"> • Limited control over the SAC structure • Heterogeneous active sites • Post/pre-treatments are required to limit the aggregation of isolated SA 	96
Atomic layer deposition	Controlled single-atom-thick-layer deposition of a gas-phase precursor on a solid support	<ul style="list-style-type: none"> • Sub-nano level thickness control • Scalable • Surface chemical selectivity • Applicable for the fabrication of SACs with a complex micropattern 	<ul style="list-style-type: none"> • Low deposition rate • Expensive 	97 and 98
Electrodeposition	Deposition of an active metal containing an electrolyte onto the solid support	<ul style="list-style-type: none"> • Scalability • Economical • Better control over the extent of the active element deposition 	<ul style="list-style-type: none"> • Possible inhomogeneous coating of SA on the host 	99
Galvanic replacement	Oxidation of one metal (support) as a sacrificial template by other metal ions (SA component) that have a higher reduction potential	<ul style="list-style-type: none"> • Better control over the morphology (size and shape, composition) of the SAC 	<ul style="list-style-type: none"> • Suitable for SA metals with a higher reduction potential than that of the host metal 	79, 100–102 and 104
Pyrolysis	Metal on the C-based host at pyrolysis temperature removes its sacrificial host template, leaving SA on the C support	<ul style="list-style-type: none"> • Robust precursors • Good thermal stability and recyclability 	<ul style="list-style-type: none"> • Requires high-temperature (>800 °C) • Applicable only for an SA on a C support • Less control over the SA loading 	34
High-temperature atomic migration	Subjecting SA species from the precursor to high-temperature so they get trapped and stabilized onto the support phase by atomically migrating under elevated temperature (≥ 800 °C)	<ul style="list-style-type: none"> • Sinter resistant • Suitable for thermocatalysis under elevated temperature conditions 	<ul style="list-style-type: none"> • Choice of the SA is restricted to metals with a low free energy • Support must be sinter resistant or must be able to undergo stable phase transition without affecting the ability to stabilize the SA 	106

Co in the lattice.³⁴ It also enabled control over the coordination numbers by varying pyrolysis temperature. A regular reduction in coordination number 4 to 2 was observed upon the increase in temperature from 800 °C to 1000 °C.

(7) High-temperature atomic migration: this is an outright method to achieve the atomic dispersion of thermally stable SACs by subjecting SA species from the precursor to get them to atomically migrate under elevated temperature (≥ 800 °C), then trapped and stabilized onto the support phase.¹⁰⁵ Generally, a high-temperature treatment will be destructive to the precursor host system. This is because the high-temperature will facilitate the migration of the SA with high free energy in

the host system, which then has the tendency to form larger aggregates, clusters, or nanoparticles (otherwise called Ostwald ripening).¹⁰⁶ Thus, one way to control the sintering possibility is to use noble metals with a low free energy in defective metal oxides, like CeO₂. However, this only works for a low level loading of SA. At higher noble metal loading, it is almost impossible to achieve atomic dispersion due to the uncontrollable formation of defects at elevated temperature. An alternative approach for achieving a higher SA loading on the supports hence can be formulated by considering the thermal behavior of the support material. For instance, Yan *et al.*¹⁰⁶ developed a sintering resistant (thermally stable) SAC by atomically



dispersing Pt onto MnO_x , taking advantage of its existence in multiple oxidation states and possible structural reconstruction during thermal treatments. During calcination at 800 °C, the initial Mn_3O_4 lattice support was reformed into Mn_2O_3 in the presence of O_2 and water vapor. In the latter case, the atomic dispersion of Pt SA was further stabilized by the strong interaction existing between the support Mn_2O_3 (Mn in a high valence state with a possibly increased number of coordination) and Pt atoms.

2. SACs for CO_2 hydrogenation

Carbon dioxide is the major greenhouse gas as its emission is leading to increased global warming and unpredictable weather change.^{108,109} The conversion of CO_2 into value-added chemicals is in great demand owing to its harmful impact on the environment and humans. However, the direct conversion of CO_2 into value-added chemicals, like methane, methanol, carbon monoxide, urea, formic acid, ethanol, ethylene, ethane, acetic acid, acetone, and propanol (Fig. 5)^{110–112} is still challenging due to the high thermodynamic stability of CO_2 . Typically, the structural characteristics of catalysts have a significant impact on the CO_2 reduction reaction process. Mainly, CO desorption from a catalyst's surface is the key step for the selective conversion of CO_2 . If CO desorption is favorable, then the reverse water gas shift reaction (RWGS) is dominant. Otherwise, additional hydrogenation occurs to produce compounds like methane or formic acid.^{113–115} The formation of CO from CO_2 requires a large amount of energy and has a high $\Delta G_{298\text{K}}^\circ$ of 28.6 kJ mol^{-1} . However, while the conversion of CO_2 to methane is thermodynamically favored ($\Delta G_{298\text{K}}^\circ = -132.4 \text{ kJ mol}^{-1}$), conversion with high selectivity toward methane at low

temperature is a real challenge and requires efforts to improve the catalytic activity at lower temperature.^{113,116–118} Additionally, CO_2 hydrogenation to methanol is also thermodynamically favorable ($\Delta G_{298\text{K}}^\circ = -10.7 \text{ kJ mol}^{-1}$) and exothermic, which means it needs to be performed at low reaction temperature. Furthermore, the formation of formic acid is comparatively less thermodynamically favorable and requires an alkaline reaction medium or additives for the reaction to proceed.^{119–121}

Recently, Wu *et al.*¹²² and Liu *et al.*⁵⁰ reported mechanistic insights into the electrocatalytic, photocatalytic, and thermocatalytic conversion of carbon dioxide into various chemicals using SACs. To scale-up the conversion of CO_2 to the industrial level, thermocatalysis is an easier approach, but it involves a high reaction temperature.¹²² The conversion of CO_2 into various value-added chemicals involves, first the formation of CH_4 , CH_3OH , HCOOH , and C_{2+} fuels or hydrocarbons and then thermocatalytic CH_4 reforming into H_2 generation (Fig. 6), making it environmentally benign and sustainable process.⁵⁹

As this review focuses on the thermocatalytic conversion of CO_2 into methane using SACs, a discussion on the general mechanism for CO_2 conversion to methane (as illustrated in Fig. 7) would be helpful. The typical CO_2 hydrogenation can be divided into two major accepted pathways: (1) CO pathway and (2) formate pathway.¹²³

As shown in Fig. 7, in the formate pathway, formate species serve as the primary intermediate products. This is referred to as the CO_2 associative pathway of the CO_2 hydrogenation reaction. In this pathway, initially chemisorbed CO_2 species can transform into bidentate formates (HCOO^*) before progressing to formic acid (HCOOH), with CH_4 produced as a result. However, in the CO pathway, known as the dissociative pathway of the CO_2 hydrogenation reaction, the adsorbed $^*\text{CO}_2$ species dissociate into $^*\text{CO}$ and $^*\text{O}$. Consequently, the generated $^*\text{CO}$ species may further dissociate into carbon species ($^*\text{C}$). These carbon species ($^*\text{C}$) afterward undergo hydrogenation to form CH_4 through interaction with dissociated H_2 molecules that remain on the metal particles and desorb from the catalyst surface. Meanwhile, $^*\text{O}$ species can react with hydrogen to generate H_2O .^{76,124,125}

3. CO_2 methanation using SACs

The conversion of CO_2 to methane is captivating as it can replace the traditional fossil fuels and can be utilized for the synthesis of syngas. Methane (CH_4) is a beneficial fossil fuel for the environment since it produces more heat and light energy per mass than other hydrocarbons, such as coal and gasoline that have been processed from oil. The hydrogenation of CO_2 at lower reaction temperature is a challenging task but is highly desirable. In this regard, various efforts have been made by researchers to enhance the selectivity toward methane rather than CO. For these reactions, the catalysts should be stable and active at high reaction pressure and temperature, and tuning of the catalysts is usually required, in which the interfacial effect between the metal atom and supported material plays a vital role. The use of a support material prevents the metal atoms or particles from agglomeration and preserves the activity of the

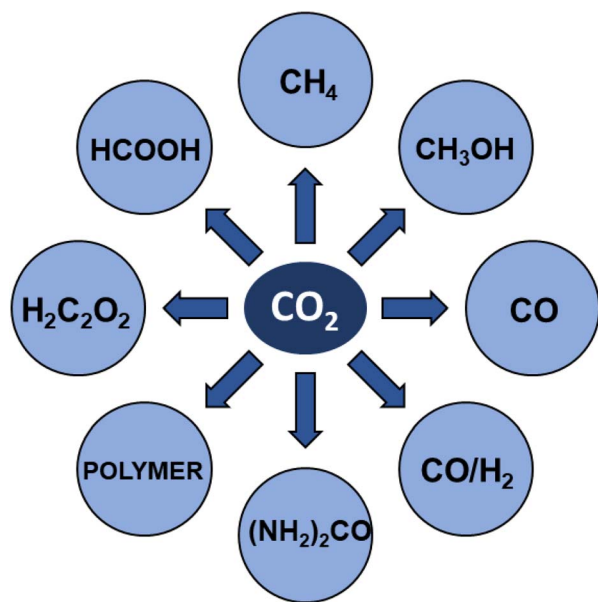


Fig. 5 Schematic illustration of the conversion of CO_2 into various value-added chemicals.



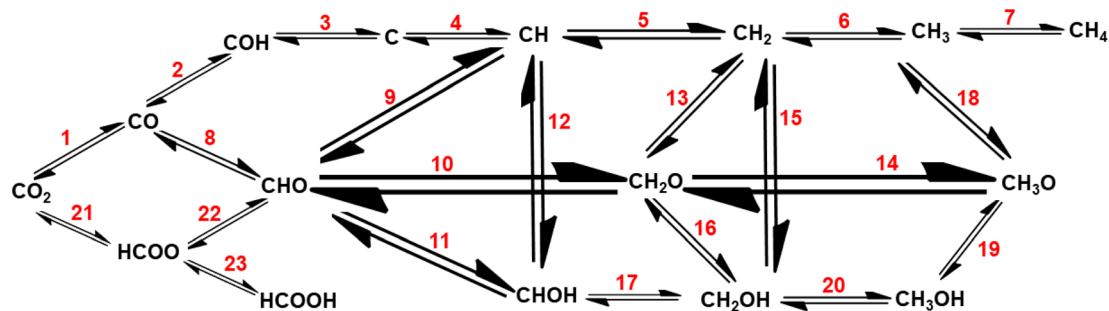


Fig. 6 Mechanistic reaction pathways for CO₂ hydrogenation to various reduced forms, like methane, methanol, and formic acid.

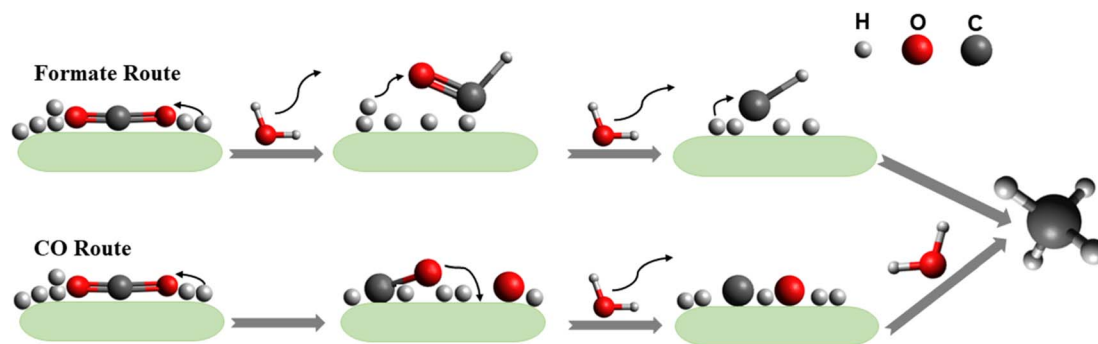


Fig. 7 General routes for CO₂ hydrogenation through the CO route and formate route.

metal component. However, the interfacial effects between the metal and support are highly affected by the particle size, surface area, porosity, acidity, and basicity of the catalyst, which consequently affects the catalytic activity and selectivity of the material for the conversion reaction. Also, the utilization of supported metal catalysts is a better alternative to deal with the issue of the costly Ru-, Pt-, and Pd-based catalysts. In such metal catalysts, a very small amount (4–6 wt%) of precious metal is doped or dispersed over a low-cost metal support in the SAC. This approach decreases the overall cost of metal catalysts without compromising the catalytic activity.^{126,127} The atomically dispersed single-atom metals provide the active sites for the reaction and maximize the catalytic performance.^{128,129} In this context, zeolite-based SACs were synthesized to produce bi-functional materials, which have high catalytic activity as well as good water sorbent capability for CO₂ methanation. Wei *et al.*¹³⁰ synthesized 13X zeolite as a support material for SACs, where CeO_x particles acted as a separator for Ni single atoms. The catalyst was synthesized by the chelation of Ni and Ce ions using citrate and was diffused over 13X zeolite. They observed that the Ce loading played a significant role in tuning of the catalyst's properties, like basicity, reducibility, and activity for the reaction. The balanced basicity and acidity allowed a well-managed interaction of CO₂ with the surface of catalyst, so that the selectivity toward methane was not compromised. Among all the synthesized catalysts, 5%Ni2.5%Ce13X showed the highest conversion and selectivity toward methane at 360 °C for 200 h timescale. Furthermore, Fan and coworkers¹³¹ synthesized a Ru-supported catalyst on porous hexagonal boron

nitride (pBN) and utilized this for CO₂ conversion into methane with high conversion and selectivity. The use of a supported material can reduce the overall cost of the catalyst without compromising its activity and thermal stability. In a Ru supported over pBN catalyst, the pBN facilitated the dispersion of Ru and generated defects *via* B and N coordination, which reduced the valence state of atomic Ru. The Ru-supported pBN catalysts showed superior catalytic activity toward methane with ~93.5% selectivity, whereas the Ru nanoparticles showed a lower conversion and selectivity during 110 h at 350 °C. Density functional theory (DFT) simulations proved that the increase in the CH₄ generation rate was due to both the atomic-scale size and the low valence state of atomic Ru supported on pBN (Fig. 8). When they increased the Ru loading, aggregation occurred, leading to the formation of Ru nanoparticles and a low dispersion of Ru over the pBN support and as a result, the CO₂ methanation selectivity decreased.

Moreover, a comparison between a plasma-decomposed catalyst and a thermally decomposed catalyst was performed for Ni over a ZrO₂ catalyst. Also, the interaction between Ni and the ZrO₂ support was studied by Jia and coworkers¹³² for the CO₂ methanation reaction, where Ni/ZrO₂ with a uniform dispersion of Ni was prepared *via* a plasma decomposition method and thermal decomposition method. The plasma-decomposed catalyst had a more uniform dispersion of Ni metal over ZrO₂ than the thermally synthesized catalyst. Owing to the high dispersion of Ni over the zirconia support, its activity was enhanced, which led to a quicker dissociative adsorption of H₂ and H-spillover in the plasma-decomposed catalyst. This H-



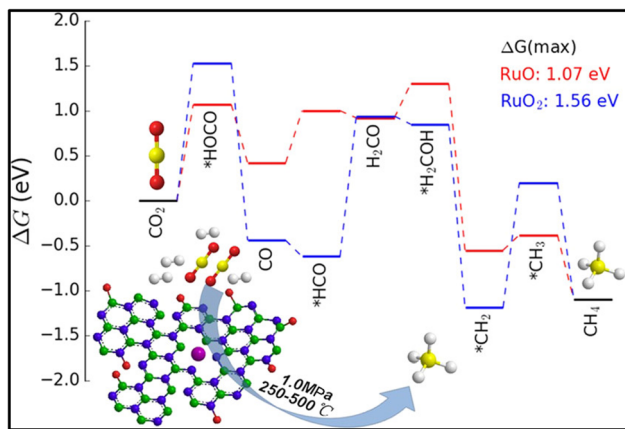


Fig. 8 Energy profile diagram for methane formation using Ru/pBN SACs. Adapted with permission from ref. 131. Copyright 2019 American Chemical Society.

spillover increased the availability of H atoms for CO₂ hydrogenation and generated oxygen vacancies on the surface of the zirconia. The oxygen vacancies ultimately increased the basicity to facilitate the CO₂ adsorption and activation capability at the surface of the catalyst. Consequently, a 71.9% conversion with 69.5% selectivity at 300 °C toward methane with a 0.61 s⁻¹ turnover frequency (TOF) was achieved at a low reaction temperature. Compared to the thermally synthesized catalyst, the TOF value was only 0.39 s⁻¹ under the same reaction conditions with a 32.9% conversion and 30.3% selectivity toward methane. DRIFT analysis revealed that the plasma-decomposed catalyst followed the CO hydrogenation pathway at the highly exposed Ni(111) lattice plane and facilitated the decomposition of CO₂ and formates into CO, as shown in Fig. 9. The thermally decomposed catalyst had a complex Ni crystal

structure and more defects, which play important roles in formate hydrogenation.

Furthermore, to stabilize Ru and Ni single atoms, Rivera-Cárcomo *et al.* synthesized CNT- and TiO₂-supported SACs and utilized them for CO₂ hydrogenation.¹³³ Both the CNT and TiO₂ supports were modified to create oxygen vacancies to stabilize the Ru and Ni single atoms. When the supported materials were utilized without creating defects, then nanoparticles and clusters of Ru and Ni were obtained. The doping of Na occurred when synthesizing TiO₂-supported SACs, due to the presence of NaBH₄. Further, the same authors synthesized Na-doped CNT for comparing the catalytic activity. Among all the synthesized catalysts compared for the CO₂ hydrogenation reaction, the Ni-based catalysts all showed less activity than the Ru-based catalysts. Surprisingly, the Na-doped TiO₂ catalyst showed enhanced activity and stability compared to the CNT-supported catalyst. However, Ni on the CNT-based SAC showed a higher selectivity toward CO formation, whereas the Ru/CNT-based SACs showed excellent selectivity toward methane formation. The charge transfer between the support and metal was the main reason for the high selectivity of the Ru SACs. They demonstrated that the electron-rich metal surface produced CH₄, while the electron-deficient metal surface produced CO more selectively. Recently, an interesting result on a single atom alloy was reported by Tu *et al.*,¹³⁴ whereby an Ir/Ni single atom alloy was synthesized *via* an impregnation method and utilized for CO₂ methanation using a homemade mechanical reactor with an attached electromotor for controlling the vibration frequency. In the Ir–Ni catalyst, Ir was atomically dispersed and coordinated with the surface Ni atoms in the form of an Ir–Ni alloy, while the reduced Ir⁰ species presented an electron-efficient state. This Ir/Ni catalyst displayed superior catalytic performance with a TOF of 10 244 h⁻¹ for methane formation and a 220 h lifetime at 350 °C without deactivation. When

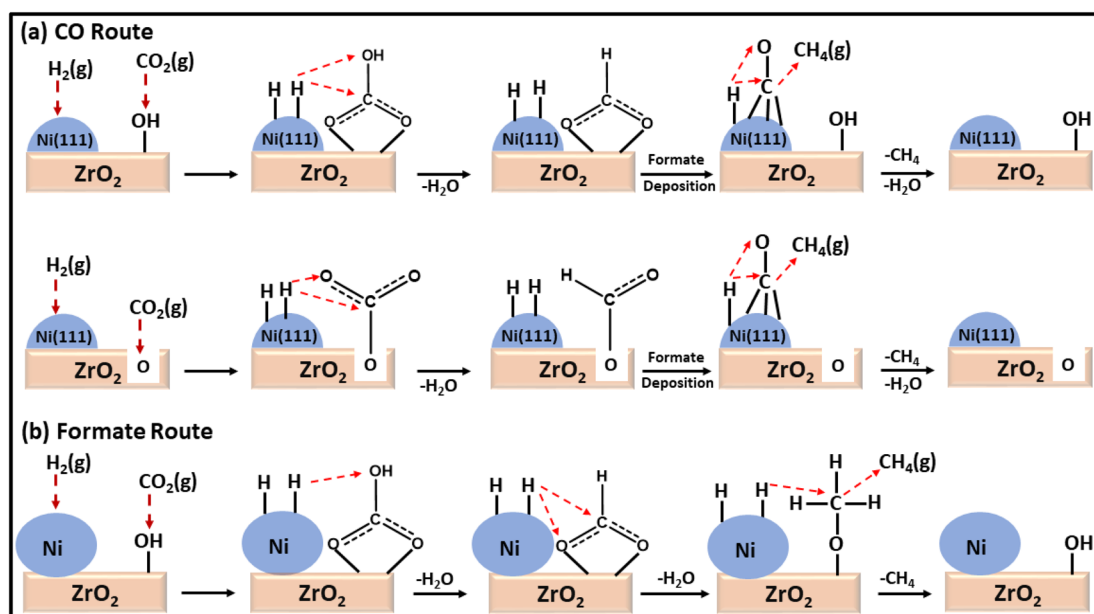


Fig. 9 Mechanistic pathways over (a) Ni/ZrO₂-plasma and (b) Ni/ZrO₂-calcined catalysts for CO₂ methanation.



compared to the Ir–Ni/Al₂O₃ catalyst supported by an oxide, the Ir–Ni catalyst demonstrated superior coke-resistant performance. The multiple oxidation states of Ni, *i.e.*, Ni⁰ and oxidative Ni²⁺, facilitated CO₂ adsorption while the Ir–Ni alloy provided the reaction site for H₂ activation and the subsequent hydrogenation reaction.

Moreover, it was reported by Liang *et al.* and Kwak *et al.* that the selectivity of the CO₂ hydrogenation reaction could be shifted from CO to methane by modifying the catalyst.^{51,135} Liang *et al.*¹³⁵ synthesized a Co single atom supported on SBA-15 *via* an ALD method, where the CoO_x thickness was controlled by the ALD cycle numbers. Based on the ALD cycles, the catalysts were named as 50Co/SBA-15 and 500Co/SBA-15. Co²⁺ was stabilized by strong Co²⁺–O–Si bonds when the catalyst was exposed to H₂ gas. The catalyst with an ultrastable Co²⁺ single atom with SBA-15 supported *via* Co–O–Si bonds exhibited 99% selectivity with a 304.6 mol CO₂/mol Co/h TOF toward CO and was stable for up to 500 h at 600 °C. In contrast, the 500Co/SBA-15 catalyst had a weak interaction bond for Co–O–Co, whereby the Co species were partially reduced to metallic Co and this metallic Co further enhanced the product selectivity toward methane. Similarly, in the 300Co/SBA-15 catalyst, the methane selectivity increased with increasing the reduction temperature due to the increased metallic Co at higher reduction temperatures. The experimental and DFT studies revealed that on the single atomic Co²⁺ catalyst, shifting between the octahedral and tetrahedral fields played an important role and the catalyst exhibited H-assisted CO₂ direct dissociation, as shown in Fig. 10. Similarly, Kwak *et al.*⁵¹ reported that a shifting of the catalytic activity could be observed from CO to methane when 3D metal clusters were formed with a higher Ru metal loading in a Ru single atom over an Al₂O₃ support. The Ru metal loading was varied from 0.1–5% and with increasing the Ru content, and the selectivity toward CO and methane was analyzed. When the metal content was low, *i.e.*, ≤0.5%, the metal was present in

the active metal phase with a high dispersion over alumina, and then the CO selectivity was high. As the metal loading of Ru increased from 1% and above, large metal clusters were formed, which led to a decrease in CO formation together with an increase in CH₄ formation. The results revealed that the size of the metal clusters decides the production selectivity in the CO₂ hydrogenation reaction. Additionally, the Arrhenius plots revealed activation energies of 82 and 62 kJ mol^{−1} for CO and methane formation, respectively, which confirmed that either the formation of methane followed a different reaction pathway without forming CO as an intermediate or by utilizing different active sites. This result was further confirmed by the difference in onset temperature between CO₂ (~325 °C) and CO (~450 °C) reduction, *i.e.*, ~125 °C difference.

Moreover, dual single-atom catalysts or dual atom catalysts (DACs) have been extensively explored due to their better performance compared to SACs.¹³⁶ Owing to the presence of heterometal sites, the catalytic activity is enhanced with a few advantages: (1) one metal atom acts as the active site while the other has an electronic regulatory role; (2) both metal centers act as active sites to enhance the catalytic activity due to a synergistic effect;^{137,138} (3) the scaling relationship limit (SRL) based on SACs is broken by the adsorption effect, in which, providing more sites alters the adsorption structures. Recently, dual metal atom catalysts have been explored for various conversion reactions, like CO₂, N₂, and O₂ gases. Consequently, Zhang *et al.*¹³⁹ explored the efficiency of a dual-active-site tandem catalyst (Ru₁Ni/CeO₂) made of Ru single atoms (Ru₁) and Ni nanoparticles. The results showed a substantially greater efficiency in the case of Ru₁Ni/CeO₂ than that of the Ru₁/CeO₂ and Ni/CeO₂ catalysts. The Ru₁Ni/CeO₂ catalysts showed a 90% CO₂ conversion and ~99% CH₄ selectivity at 325 °C. DFT calculations and the experimental results confirmed the synergistic effect between the Ru₁ and Ni sites, which boosted the overall methanation pathway (Fig. 11). As shown in Fig. 11, first

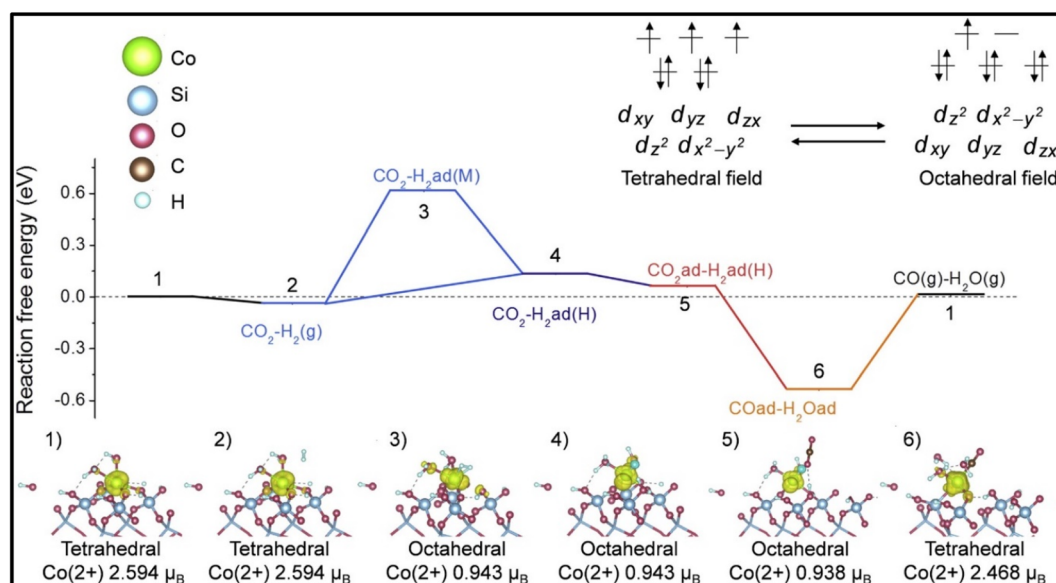


Fig. 10 Mechanistic pathway over Co/SBA-15 catalyst for methanation. Adapted with permission from ref. 135. Copyright 2024 Elsevier B.V.



the dissociation of CO₂ goes on at the Ru single atom, which is further hydrogenated to produce *CHO over the Ni NPs. Afterward, this *CHO is converted into methane over Ni sites of the Ni NPs. Hence, the Ru single atom species help to convert CO₂ to CO, whereas the Ni sites promote the reaction further from CO to CH₄.

In another report,¹⁴⁰ a Ru₁Ni SAA catalyst was synthesized over a SiO₂ support and named as Ru₁Ni/SiO₂. First, Ni phyllosilicate (Ni-ph) was reduced to derive Ni NPs *via* a hydrothermal preparation method and then the GRR was performed in between ruthenium chloride and the as-synthesized Ni NPs. To compare the catalytic activity, a methanation test was performed, where the Ru₁Ni/SiO₂ DAC showed higher catalytic activity than Ni/SiO₂ and other previously reported Ni-based catalysts. The DAC showed a TOF of $40.00 \times 10^{-3} \text{ s}^{-1}$, which was significantly higher than that of Ni/SiO₂, *i.e.*, $4.40 \times 10^{-3} \text{ s}^{-1}$. Based on DFT calculations and the experimental results, the reaction mechanism was proposed and is shown in Fig. 12. It was mentioned that initially, over the Ru and Ni sites of DAC, the adsorption of CO₂ occurred. Consequently, this CO₂ molecule dissociated into CO*, which further hydrogenated into methane over the Ru single atom. The atomically dispersed Ru species on the Ni surface exhibited a pronounced interaction with nickel atoms, resulting in significant charge transfer from Ru atoms to Ni atoms, facilitating the dissociation of CO₂ into CO* for the high catalytic activity and selectivity toward methane formation.

In addition, Kikkawa *et al.*^{141,142} reported a Ni–Pt alloy (iso-Pt) produced on aluminum oxide (Al₂O₃) *via* an impregnation method under the pretreatment of hydrogen for reduction. The Ni–Pt alloy over alumina (Ni₉₅Pt₅/Al₂O₃) had isolated Pt atoms surrounded by Ni atoms, and exhibited a relatively high CH₄-formation rate owing to a lower activation energy than that of the bulk-Ni (Ni/Al₂O₃) and bulk-Pt (Pt/Al₂O₃) catalysts. It was discussed that the iso-Pt species play a dual role: one is for H₂ dissociation at low reaction temperature and the other one is

tuning the d-electronic state by alloying with Ni atoms. The iso-Pt atoms enhanced the ability of the nearby Ni atoms to encourage more methanation with hydrogen, while weakening the C–O bond of the CO species connected to the Pt atoms. Through the electrical interactions between Pt^{δ-} atoms and the surrounding Ni^{δ+} atoms, isolated Pt atoms were produced on the surface of the Ni₉₅Pt₅ solid solution alloy, as shown in Fig. 13. Additionally, they also compared the catalytic activity of Ni₉₅Pt₅/Al₂O₃ with Ni/Al₂O₃. Moreover, they extended the study into the cooperative effect between the neighboring Ni atoms and isolated Pt atoms in the Pt–Ni alloy using *in situ* and transient Fourier transform infrared (FT-IR) studies and kinetic studies for the production of methane from CO₂. They observed that the formation of CO in the bridge form as an intermediate between the iso-Pt and Ni atoms played a significant role in selective methane production. The transient variations in the FT-IR spectra showed that the bridging CO species hydrogenated the CO species connected to the isolated Pt atoms in the direction of CH₄. The kinetic studies revealed that after the addition of Pt species in the Ni catalyst, the H₂-dissociation step became the rate determining step instead of the hydrogenation of the surface carbon species step. Also, the carbon species on the Ni–Pt alloy were hydrogenated more quickly due to the strong H₂-dissociation capacity of the Pt atoms. These results confirmed that the isolated Pt atoms of the Ni–Pt catalyst play a dual role, acting as effective H₂ dissociation sites to encourage hydrogenation and as adsorption sites for the on-top CO species to afford a high selectivity toward CH₄ (Fig. 14).

In addition, Kwak *et al.*¹⁴³ synthesized Pd + La₂O₃/MWCNT, a bimetallic catalytic system combining SACs to boost the catalytic activity. Here La₂O₃ acted as a promoter, while atomically dispersed metallic Pd facilitated the dissociation of H₂, and the size of the metal particles determined the selectivity of the product of the CO₂ hydrogenation. With this approach, atomically dispersed Pd-based catalysts supported on Al₂O₃ and

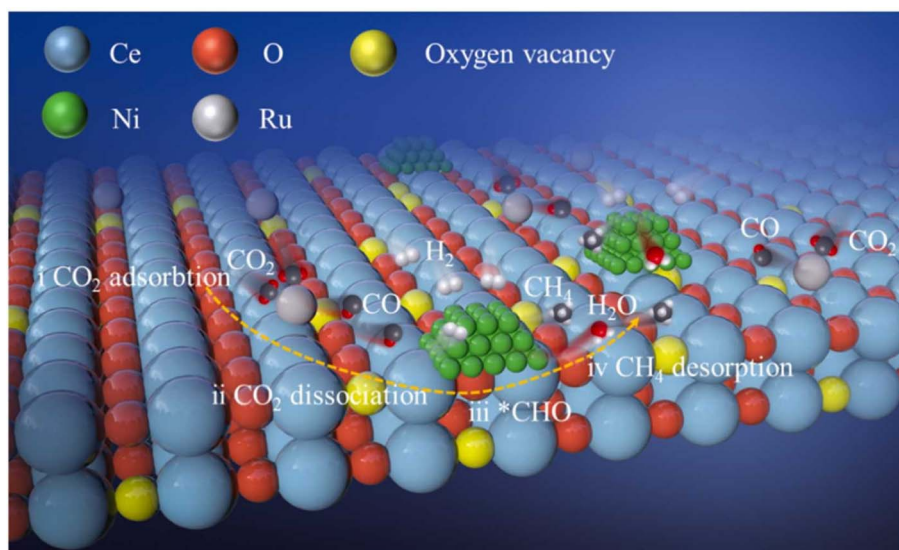


Fig. 11 Schematic representation of CO₂ methanation using Ru₁Ni/CeO₂ DACs. Adapted with permission from ref. 139. Copyright 2023 Elsevier B.V.



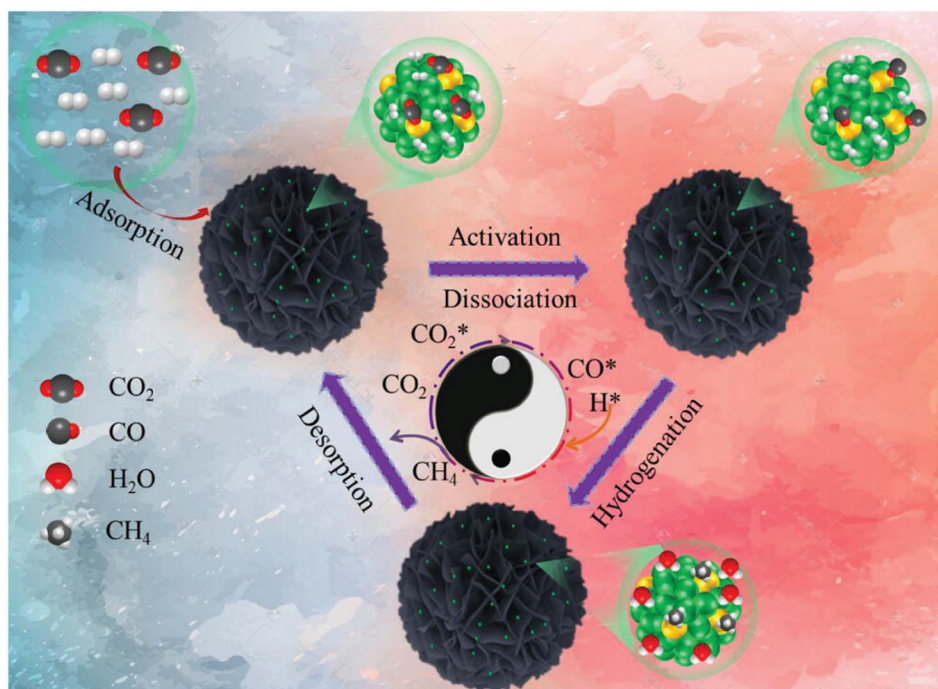


Fig. 12 Schematic representation for the production of CH₄ over Ru₁Ni/SiO₂. Adapted with permission from ref. 140. Copyright 2023 Wiley-VCH GmbH.

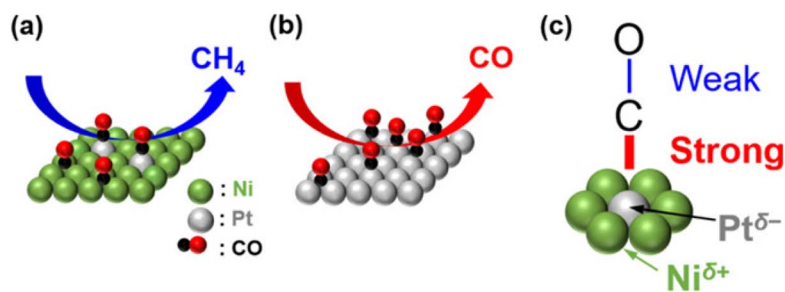
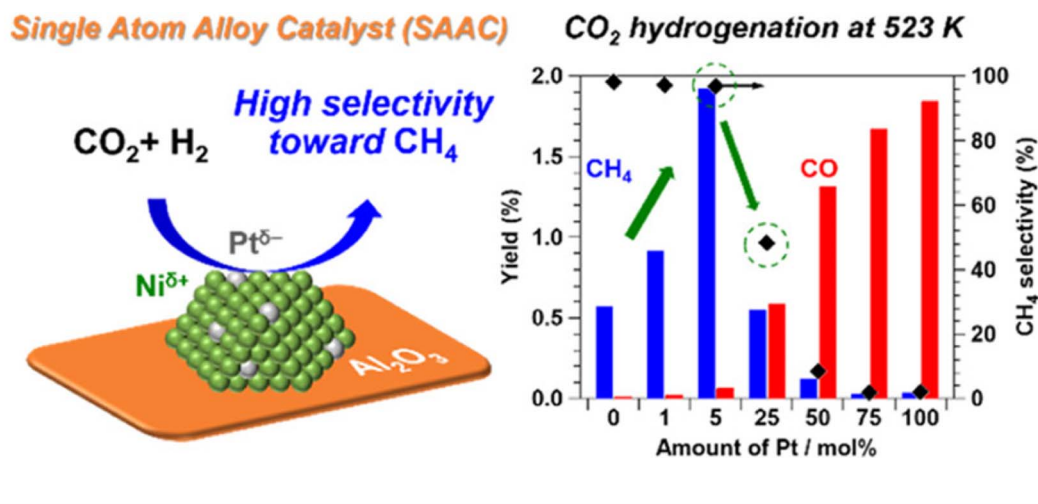


Fig. 13 Representation of the adsorbed CO species on the surfaces of the (a) iso-Pt catalyst and (b) bulk-Pt catalyst, and (c) the adsorbed CO model on a single Pt^{δ-} atom encircled by Ni^{δ+} atoms. Adapted with permission from ref. 141. Copyright 2019 American Chemical Society.



Bifunctional roles of isolated Pt atoms in Pt–Ni single atom alloy catalyst (SAAC)

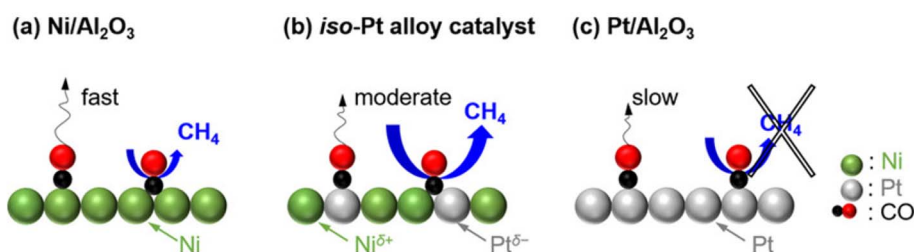
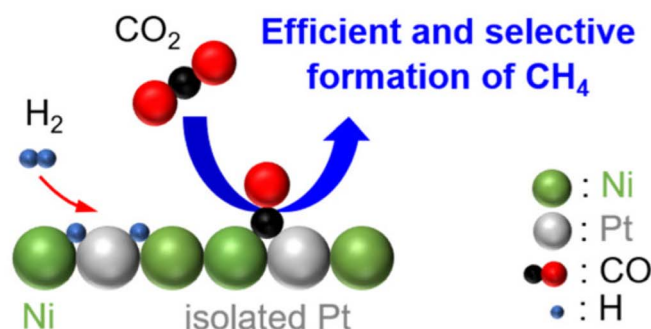


Fig. 14 Graphical representation of the adsorbed species and CO₂ hydrogenation over (a) Ni/Al₂O₃, (b) the iso-Pt catalyst, and (c) Pt/Al₂O₃. Adapted with permission from ref. 142. Copyright 2020 American Chemical Society.

Pd over multiwalled carbon nanotube (MWCNT) catalysts were also synthesized. Initially, when their activity for the CO₂ hydrogenation reaction was analyzed, the Pd/Al₂O₃ SACs showed high catalytic activity, while Pd/MWCNT was completely inactive for the hydrogenation of CO₂. However, when La₂O₃ oxide was introduced to the inactive Pd/MWCNT catalyst, then the catalyst showed superior catalytic activity compared to both the Pd/Al₂O₃ and Pd/MWCNT catalysts. Nonetheless, selectivity and conversion toward methane were only obtained in the case of Pd/Al₂O₃, as shown in Fig. 15. In this catalyst, large Pd clusters were present over an alumina support, on which the initially generated CO was readily hydrogenated to methane, while the selectivity toward methane increased as the temperature increased due to sintering of the Pd active metal phase.

Moreover, Shin *et al.*¹⁴⁴ synthesized an atomically dispersed Pt catalyst over barium zirconate (BaZrO₃ = BZ) perovskite with Co nanoparticles and compared the activity with γ -Al₂O₃-supported catalysts. Generally, the catalyst must transport cooperative hydrogen between the Pt metal support and Co species to produce the desired functionality, so that the dissociation of CO₂ occurs on cobalt and that of H₂ on the Pt metal. In this regard, they synthesized atomically dispersed Pt on the surface of Co nanoparticles. The catalyst was synthesized by the immobilization of Pt particles over the Co/BZ support with 1 wt% Co and 0.2 wt% Pt loading. They observed more than a 6-fold increase in CH₄ formation rate with Pt on the Co/BZ support compared to that with the alumina support at 325 °C. This enhancement in activity in the case of the BZ support was due to the strong interaction between the Co particles and the

BZ support, and the dissociation of CO₂ and H₂ was thus facilitated on the surface of Co and Pt, respectively. Despite this, more than 70% CH₄ selectivity was obtained, which remained constant throughout the measured temperature range of 250–350 °C. The selectivity was 80% even at 325 °C, compared to only 43% selectivity with the γ -Al₂O₃ support. Additionally, when yttria was doped on zirconia (5 and 30 atom%, respectively), a decrease in catalytic activity was observed compared to when yttria was supported by BZ. However, even at low substitution levels, the activity was still higher than that of the γ -Al₂O₃ support.

In contrast, there are several reports where nanoparticles and nanoclusters have shown higher selectivity toward methane over SACs. However, isolated active sites of SACs suppress the side reactions and generate products with high selectivity. In this regard, the metal particle size must be taken into consideration, as it plays a significant role in the distribution of active sites and consequently in the reactivity and selectivity. The size effect in catalysis is a highly disputed and intricate subject, primarily because the most effective sizes of metal particles can vary significantly across different catalytic setups.⁶⁰ It often requires careful experimental optimization and theoretical modeling to understand the interplay between the particle size, catalytic activity, and reaction mechanisms. Guo *et al.*⁶⁰ studied the effect of the size of the particles in supported SACs for the CO₂ hydrogenation reaction to methane in detail. They synthesized a Ru single atom supported on CeO₂ nanowire, Ru nanoclusters (1.2 nm), and Ru nanoparticles (4 nm) and utilized these for CO₂ hydrogenation to methane. It was observed that



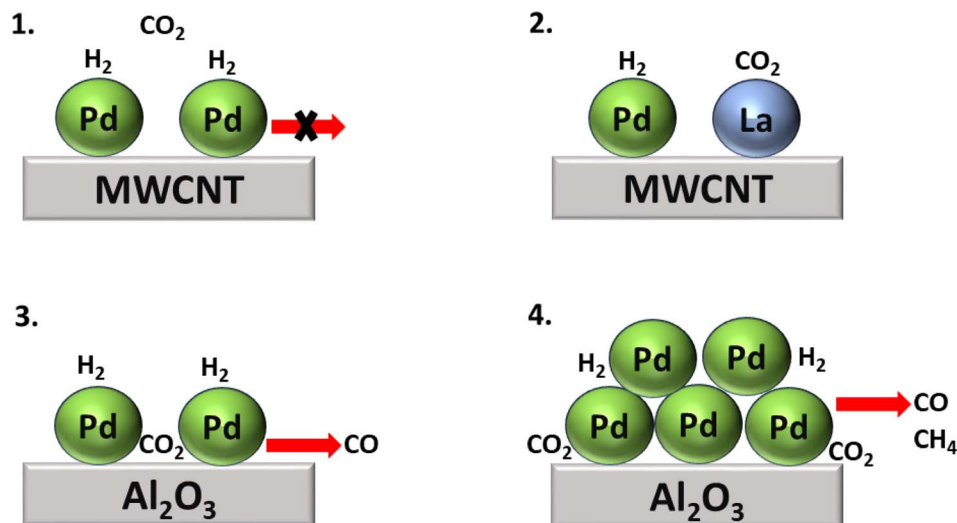


Fig. 15 Comparative study of CO₂ hydrogenation over (1) Pd/MWCNT, (2) Pd + La₂O₃/MWCNT, (3) large Pd cluster over Al₂O₃, and (4) Pd/Al₂O₃ SACs.

with the increase in the size range, the selectivity and activity toward methanation varied. The TOFs for methane selectivity with the Ru/CeO₂ SACs, Ru nanoclusters, and Ru nanoparticles were 4.59×10^{-3} , 7.41×10^{-3} , and $5.30 \times 10^{-4} \text{ s}^{-1}$, respectively at 190 °C. The highest activity and selectivity toward methanation were observed in the case of the Ru nanoclusters. *In situ* DRIFT analysis revealed that the CO route was the predominant reaction pathway with Ce³⁺-OH sites and Ru sites near the metal-support interfaces acting as the active sites for CO₂ dissociation and carbonyl hydrogenation, respectively. DFT calculations revealed that as the strong metal-support interaction (SMSI) of CeO₂ with Ru atom decreased, CO adsorption on the surface of the catalysts increased, whereas weakening of the

H-spillover effect enhanced the removal of water. Consequently, the activity and selectivity for methanation were greatly enhanced in the case of the Ru nanoclusters at low reaction temperatures, as shown in Fig. 16. The SMSI and H-spillover play important roles that can affect the activity of the catalysts with the size regimes, which was balanced in the case of the Ru nanocluster supported on CeO₂ support.

Furthermore, Matsubu *et al.*¹⁴⁵ compared the catalytic activity of Rh nanoparticles and Rh supported on TiO₂ SACs for CO₂ hydrogenation reactions. They performed diffuse reflectance infrared Fourier transform spectroscopy (DRIFTS) analysis to quantify the Rh active sites in the synthesized catalysts. There was significant relation observed between the TOF of

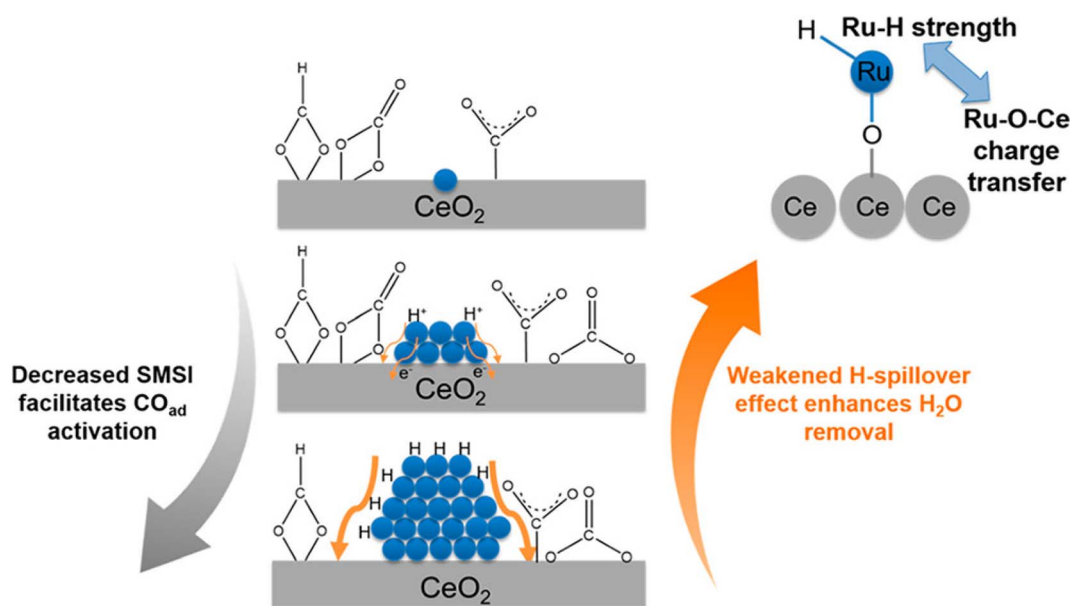


Fig. 16 Effect of the competition of SMSI and H-spillover on CO activation. Adapted with permission from ref. 60. Copyright 2018 American Chemical Society.



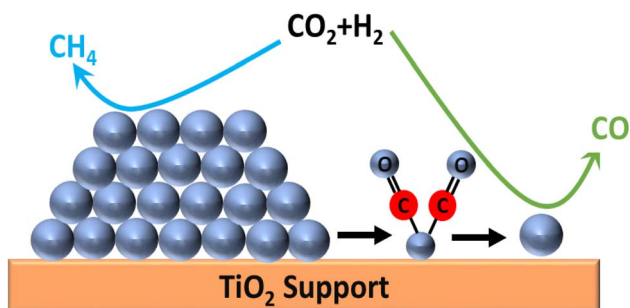


Fig. 17 CO₂ hydrogenation selectivity over Rh NPs and Rh/TiO₂ SACs.

RWGS with Rh_{iso} (Rh SACs) sites and the TOF for methanation with Rh nanoparticles (Rh_{NP}). When the CO₂ hydrogenation reaction was performed, the nanoparticles showed selectivity toward CH₄ and the SACs showed selectivity toward CO. They demonstrated that all the Rh sites in the Rh nanoparticles were surrounded by other Rh sites, which were more active for H₂ dissociation and bound CO very strongly. Consequently, sequential hydrogenation occurred, and methane formation was favored over CO desorption. However, in the case of the SACs, *i.e.*, the isolated Rh sites over the TiO₂ support, the CO bonding was weak and the selectivity toward methane decreased. They concluded that the SACs and nanoparticles supported on the same metal oxide, *i.e.*, TiO₂, followed uniquely different pathways for CO₂ hydrogenation and showed different

selectivities, where disintegration of the nanoparticles played a considerable role in controlling the stability (Fig. 17).

Additionally, Jimenez *et al.*¹⁴⁶ compared the catalytic activity of Co/SiO₂ single-atom catalysts with Co clusters for the CO₂ hydrogenation reaction. They observed that atomically dispersed Co over the SiO₂ catalyst where the isolated Co was present in the tetrahedral Co²⁺ coordination showed the highest conversion toward CO, whereas cobalt clusters (sized 2–30 nm) exhibited selectivity toward CH₄ formation under similar reaction conditions. They found that the local coordination environment of the Co metal played an important role to tune the selectivity toward methane. SACs with a distribution of Co⁰/(Co²⁺+Co⁰) of 0.122–0.363 exhibited CO₂ conversion at a 600 °C reduction temperature, while nanoparticles sized 2–30 nm exhibited CO₂ conversion at higher temperature. When the Co⁰/(Co²⁺+Co⁰) ratio was 0.363, the catalytic activity and selectivity toward methane were high whereas, then as the ratio decreases, the selectivity and conversion toward methane also decreased. The superior catalytic activity at 0.363 was estimated to be due to a synergistic effect between the metallic Co and Co²⁺. This result indicated the ensemble effect for transition metals in the subnanometer scale and the effect of the particle size for surface-sensitive reactions. Similarly, Wang *et al.*¹⁴⁷ reported Pt/CeO₂ SACs for the CO₂ hydrogenation reaction and compared their catalytic activity with nanoclustered Pt catalysts. Atomically dispersed Pt over the CeO₂ catalyst had only a 0.05 wt% loading of Pt, whereas the Pt/CeO₂ nanoclustered catalyst had

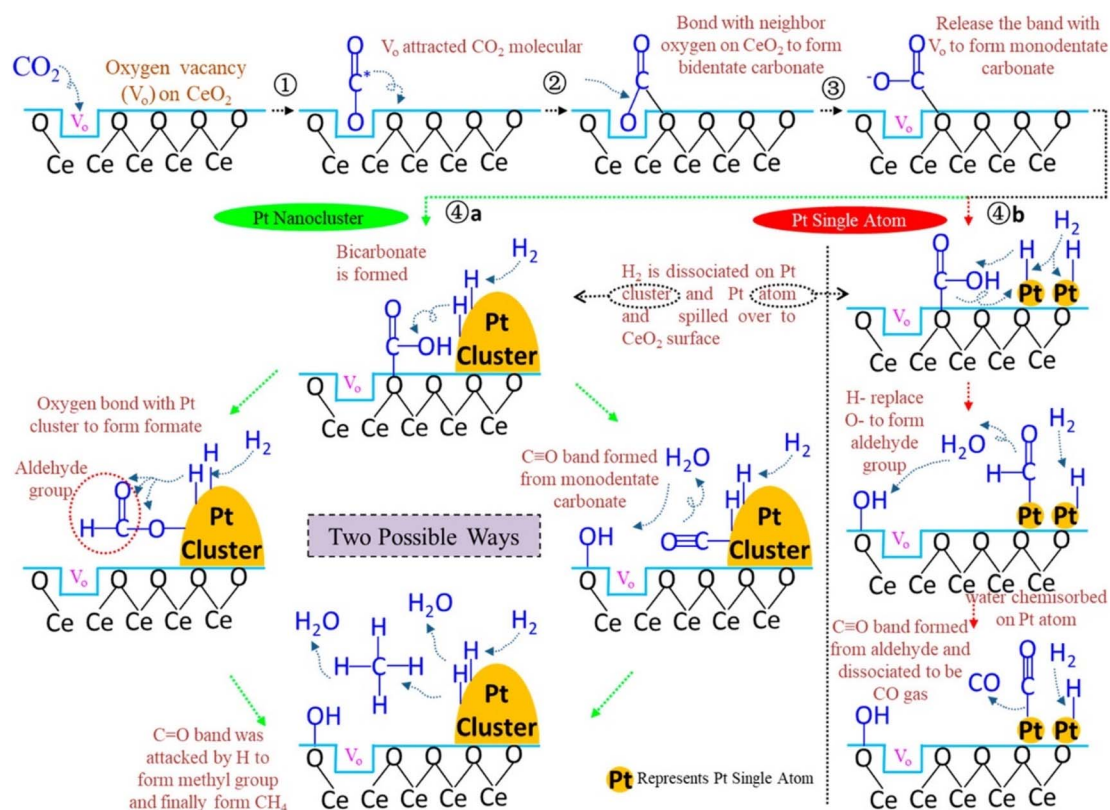


Fig. 18 Plausible reaction mechanism over the Pt nanocluster and Pt SACs catalysts. Adapted with permission from ref. 147. Copyright 2018 American Chemical Society.



a 2 wt% Pt loading. To reduce the total cost of the catalyst, the usage of noble metal should be reduced and, in this context, a significantly lower amount of Pt was utilized in these catalysts. When the reaction was performed for CO₂ hydrogenation, the atomically dispersed Pt/CeO₂ displayed a 7.2 times higher reaction rate over the nanoclustered Pt/CeO₂ catalyst even with a 40 times lower Pt loading. It was revealed that the activation of CO₂ occurred on the oxide support while H₂ dissociation occurred on the Pt metal. However, the atomically dispersed Pt SACs showed selectivity toward CO formation, whereas the nanoclustered Pt catalyst showed selectivity toward CH₄ formation. It was demonstrated that on the dispersed Pt surface, CO binds weakly, which restricts its further hydrogenation to methane and consequently the CO selectivity increases. In both the nanoclustered and SACs catalysts, the CeO₂ present is rich with surface oxygen defects where the adsorption of CO₂ occurs. As shown in Fig. 18, CO₂ molecules first interact with the oxygen defects of CeO₂ and bind with the oxygen of CO₂ in a bidentate manner to yield carbonate species. Further, with increasing the reaction temperature, this oxygen is released from the oxygen defects and generates monodentate carbonate species, and consequently at higher reaction temperatures, methane production occurs. Moreover, another mechanism pathway was also proposed where a monodentate system is first generated on the CeO₂ surface and reacts with the adsorbed H, which was observed in the *in situ* DRIFT analysis. Further, the OH⁻ of bicarbonate is replaced with adsorbed H, which generates CO *via* the cleavage of C–H bonds of the aldehyde group on the single atomic Pt of the atomically dispersed Pt/CeO₂ catalyst. This aldehyde further will not be hydrogenated to methane. This might be due to the strong bonding of the OH group with the CeO₂ support of the SAC, which reduces the H-spillover, thus hindering CO dissociation and methane formation.

4. Conclusions and future outlooks

A survey of the available reports on SACs for the hydrogenation of CO₂ highlights that the scope grows rapidly toward commercializing natural gas to a promising regime. The invention of SACs on the whole greatly alleviates the obstacle of scaling-up processes that previously used precious metals as catalysts that are highly selective and effective toward CO₂ methanation. The most comprehensive progress report on SACs attempted so far is provided with a special emphasis on the hydrogenation of CO₂ to methane gas. Based on the survey, the below conclusive points are made and further future directions are suggested:

(1) The scope for the methanation of CO₂ by SACs is still in its infancy as few systems have yet managed to achieve sufficient selectivity. This is because theoretical studies say that conventional SACs show feasibility toward the hydrogenation of CO₂ to formic acid. The bottleneck is the need to formulate SACs based on a clear understanding of the structural features that can control the product selectivity toward methanation during CO₂ hydrogenation.

(2) Elaborated data technology is required to unravel the exact structures of SACs and the complex reaction network by theoretical analysis. The state-of-the-art computational details are currently unsatisfactory in this respect.

(3) The understanding of the complex mechanisms underlying catalysis in SAC systems has far to go. It is hard to establish a general reaction mechanism for CO₂ conversion by SACs due to the limited understanding of the key reaction intermediates and the complex interactions between different phases of the SACs and reactants, even in the case of SACs with a well-defined structure.

(4) It is essential to reveal the temporal changes in the structural and electronic features of SACs under real-time reaction conditions. Exploiting new characterization tools may allow handling of these issues in the future. Many of the existing analysis tools are not sufficient to fully elucidate the complex processes. For example, the widely used *in situ* characterization tools, like *Operando* FT-IR analysis, do not yield information about weak interactions. Instead, *Operando* techniques, like near ambient pressure X-ray photoelectron spectroscopy (NAP-XPS), endorsed by low energy electron diffraction (LEED) and scanning tunneling microscopy (STM) measurements can give more in-depth information about the transient intermediates formed during CO₂ methanation.¹¹³

(5) The possibilities of providing a proper coordination environment with distinct electronic features for other low-cost metals, like Ni in SACs, which otherwise are not so promising for the methanation of CO₂, would also expand the choice of active SAs in SACs to a wider margin in the periodic table.

Conflicts of interest

There are no conflicts to declare.

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