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Green energy driven methane conversion under mild conditions

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Methane is a critical energy resource but also a potent greenhouse gas, significantly contributing to global warming. To mitigate the negative effect of methane, it is meaningful to explore an effective methane conversion process motivated with green energy such as green electricity and sunlight. The selectivity and production rate are the key criteria in methane conversion. This review provides a comprehensive overview of recent efforts and understanding in methane conversion to valuable products, including oxygenates and hydrocarbons, by taking advantage of electrocatalysis and photocatalysis. The review begins with a general understanding of C–H bond activation mechanisms. It then focuses on electrocatalytic methane conversion (EMC) with an emphasis on catalyst design for oxygenate production, and photocatalytic methane conversion (PMC) with a particular focus on hydrocarbon production, especially ethylene (C_2H_4), due to the differences in oxygen sources between the two systems. An in-depth understanding of EMC and PMC mechanisms is also discussed to provide insights for improved catalyst design aimed at selective product generation. Finally, successful catalyst designs for EMC and PMC are summarized to identify challenges in achieving highly efficient and selective production of value-added chemicals and to offer clear guidance for future research efforts in green methane conversion.

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Broader context

Methane's greenhouse effect has been shadowed by its significance in powering our society. With the carbon tax being implemented globally, methane emission becomes an emerging issue. The advances in solar energy utilization, such as solar-to-electricity and solar-to-chemical conversions, provide a green and sustainable pathway to address the methane emission issue by converting methane into valuable products under mild conditions. In order to achieve good selectivity and high production in methane conversion, different methods are applied, including electrocatalytic methane conversion (EMC) and photocatalytic methane conversion (PMC). This review will summarize the recent progress in EMC for oxygenate generation and PMC for hydrocarbon generation with the understanding about the C–H bond activation. This effort aims to shed light on innovations in methane management through solar energy.

1 Introduction

Methane (CH_4) has significantly contributed to global warming since the Industrial Revolution, with a global warming potential 84 times greater than that generated by carbon dioxide over 20 years and 28 times greater over a century.^{1,2} In 2021, global methane concentrations reached 1896.7 ppb, 2.62 times higher than pre-industrial levels (Fig. 1a).³ Currently, 580 Mt of methane is released into the atmosphere annually, with 60% of these emissions being anthropogenic. Agriculture accounts

for 24% of total emissions, followed by fossil fuel utilization at 23%, largely from the generation of associated natural gas (Fig. 1b).^{1,2,4} Flaring methane during oil production alone releases around 140 billion cubic meters of associated natural gas each year, equivalent to approximately 3.5% of the annual natural gas production in 2020.^{5,6} With growing pressure to reduce carbon emissions and the commitment to achieve net zero scenario, there is an urgent need to mitigate the methane emission issue.

Besides its environmental impact, methane is also an important industrial C_1 building block for many chemicals.⁷ Current industrial processes for methane conversion require two major steps with intensive energy consumption and carbon footprint. Generally, steam methane reforming is applied to produce syngas, a mixture of hydrogen and carbon monoxide,⁷

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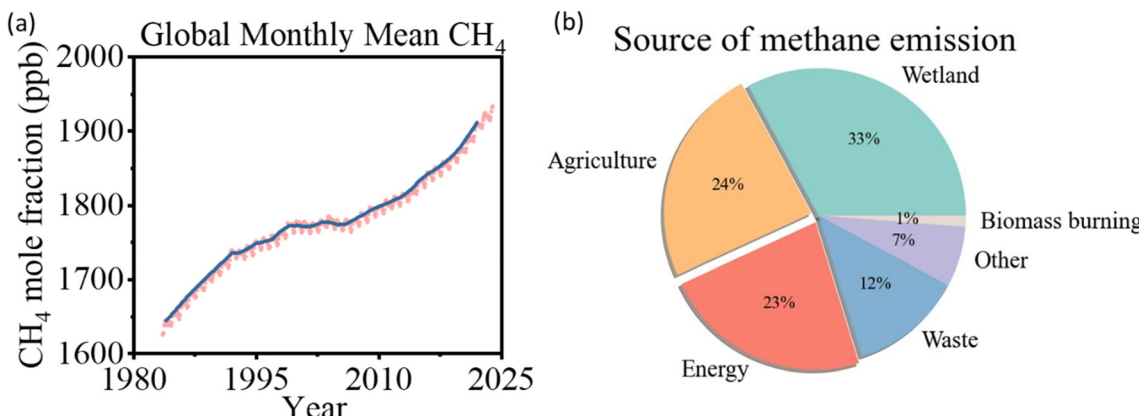


Fig. 1 (a) Global averaged atmospheric methane, data from the National Oceanic and Atmospheric Administration, 2022.³ (b) Source of methane emission, data from the International Energy Agency, 2022.¹

for many synthesis processes including Fischer-Tropsch synthesis for hydrocarbons, and Oxo-process for oxygenates.⁸ The conversion of methane into syngas is a highly endothermic reaction ($\Delta H = +206.2 \text{ kJ mol}^{-1}$), which makes the reaction process require high thermal energy input and maintain a temperature above 700 °C.⁷ In pursuing sustainable society development and green chemical engineering, it is significant to explore direct conversions of methane into desirable products *via* a mild process.

Besides thermal energy, the photons and electrons can also activate methane as shown in Fig. 2a, emerging as increasingly popular fields of research since the beginning of this millennium (Fig. 2b).^{9–11} In a thermocatalytic process, heat input is to

activate the catalyst surface and methane molecules, but the low thermal utilization efficiency significantly limits the overall conversion process. In comparison, electricity and solar energy are high-grade energy which can be used as the driving force to overcome the methane activation barrier, therefore the electrocatalytic and photocatalytic methane conversion can be effectively processed at room temperature. In particular, electrocatalytic methane conversion (EMC) studies focus on the partial oxidation of methane for oxygenates due to the aqueous environment, whereas most photocatalytic methane conversion (PMC) can achieve methane coupling for high-carbon product generation.^{12–14}

In the presence of oxygen, methane conversion will undergo a downhill reaction pathway with a negative enthalpy under

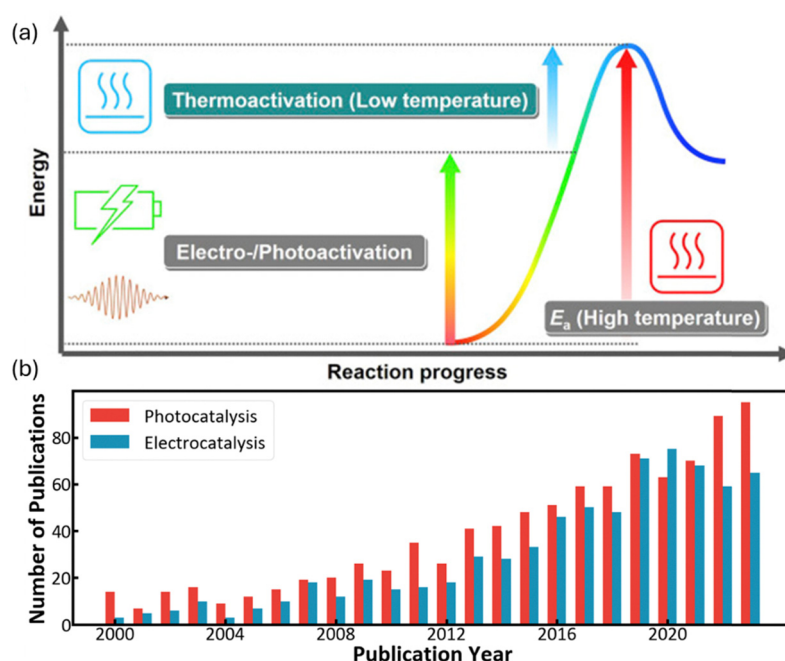
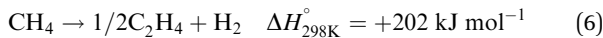
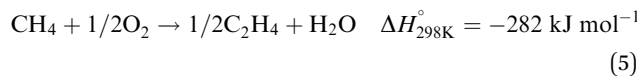
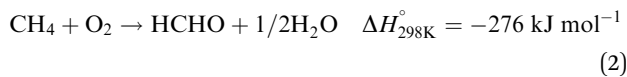


Fig. 2 (a) Energy diagram for low-temperature methane conversion through thermal-, electro- and photocatalysis. Reproduced with permission.⁷ Copyright 2019, Elsevier. (b) Number of publications that include "Methane" and "Electro"/"Photo" according to the Web of Science (as on 30 August 2024).



standard conditions (ΔH_{298} , reactions (1)–(5)).⁷ Particularly for methanol production *via* reaction (1), it can be achieved *via* a simple one-step reaction with a ΔH of -126 kJ mol^{-1} .¹⁵ Compared to the conventional coal-based methanol production process, the methane partial oxidation represents a tidy and atomic economic reaction pathway. However, these oxygenated products are hard to preserve in the reaction environment as overoxidation of methane into carbon monoxide or carbon dioxide is much more favourable as shown in reactions (3) and (4).¹⁶ Methane coupling can also be expected *via* reaction (5) through the oxidative coupling of methane (OCM). Meanwhile, methane coupling can also be achieved in a non-oxidative procedure (NOCM), but with significantly higher ΔH as demonstrated in reaction (6).¹⁷ As a comparison between the OCM and NOCM processes, the OCM shows a faster reaction rate but at the risk of deep oxidation of the products, while NOCM gives much higher selectivity but at the cost of a slow reaction rate and coke formation.^{7,17,18}



The key design of direct methane conversion is to dissociate C–H bonds of methane under mild conditions and mediate the oxidation in a controllable manner towards the desired products. In an aqueous environment, the water, on the one hand, will prevent methane from accessing the electrode surface, and on the other hand, it will be oxidized on the electrode surface for active oxygen species generation to activate C–H bonds. Therefore, the EMC process needs to balance active oxygen species generation and the oxygen evolution reaction, which has been less reviewed.^{7,15,16,19–22} As a comparison, the PMC process is generally processed in the gaseous phase, which will ensure the easy access of methane molecules and oxidants (generally O_2). Therefore, the rate limiting step is not the oxidative species generation step, but the methane activation process. By precisely controlling the methane activation step, not only ethane (C_2H_6) can be produced,^{11,23,24} but also some more valuable products such as ethylene (C_2H_4) can be produced.²⁵ For progressing research in C_2H_4 generation *via* PMC, an overview of the progress of photocatalytic CH_4 to C_2H_4 generation is necessary.

In this review, we focus on the cutting-edge advancements in methane conversion under mild conditions, particularly through EMC and PMC. The discussion includes the activation of the C–H

bond and the various reaction pathways. By comprehensively revisiting catalyst design for EMC and PMC processes, we illustrate how the oxygen source influences product selectivity, and the challenges associated with improving production rates and selectivity. Towards the conclusion, we provide an outlook on these advancements and outline potential research directions for further development in both EMC and PMC.

2 Methane C–H bond dissociation

Methane has a very stable, highly symmetric tetrahedron structure, which makes it have not only low polarization but also extremely high bonding energy of 439 kJ mol^{-1} .⁷ These features lead to harsh kinetics with low yield and poor selectivity for direct methane conversion. The biggest challenge of methane conversion lies in dissociating the C–H bonds of methane molecules, which is regarded as the holy grail of chemical reactions. Especially after the first C–H bond dissociation, the symmetry of CH_4 will be broken down, which will trigger the following C–H bond activation. The dissociation pathways can be classified into deprotonation and dehydrogenation (Fig. 3) according to the formal oxidation state (FOS) of the central carbon.²⁶

The dehydrogenation process refers to the detachment of one hydrogen atom (that is one proton with one electron) from methane, resulting in the formation of a methyl radical ($\bullet\text{CH}_3$).²⁹ Oxygen can be applied as the hydrogen acceptor to form a hydroxide radical ($\bullet\text{OH}$). Deprotonation of methane is the removal of a proton and generation of a methenium ion (CH_3^-). It will result in the carbon maintaining its FOS at $-IV$ with a tetrahedral geometry with sp^3 hybridization, which will coordinate with the active site to form a metal–carbon σ bond as shown in Fig. 3.^{26,30–32} In contrast to the deprotonation, methyl radicals generated from the dehydrogenation mechanism result in the central carbon's FOS changing from $-IV$ to $-III$ with a trigonal-like sp^2 hybridisation. It has weak interaction with the catalyst active sites as shown in Fig. 3.²⁶ From the above discussion, deprotonation of methane is an acid–base reaction due to the proton participation, which is in line with the observation that the deprotonation process usually takes place in highly polar solvents such as water and high concentration acids within homogeneous systems.³⁰

To distinguish these two mechanisms, the reaction energy profile by two pathways needs to be compared which can be clarified by the active oxygen species involved in the reaction. The active oxygen species can be electrophilic (e.g., O^- , O_2^-) or nucleophilic (e.g., O^{2-}).³³ The dehydrogenation of methane involves electron-deficient oxygen species including O^- , O_2^- from strong oxidants such as O_2 and N_2O , while methane C–H bond deprotonation generally involves with electro-saturated oxygen species (O^{2-}) as proton acceptors.³⁴ As a result, dehydrogenation is usually accompanied by strong oxidizing catalysts such as metal oxides and metal oxyhydroxides, and deprotonation requires metal or metal complexes with low oxidation states together with proton acceptors.³⁴

After the first C–H bond dissociation in methane, the remaining C–H bonds become vulnerable to further dissociation



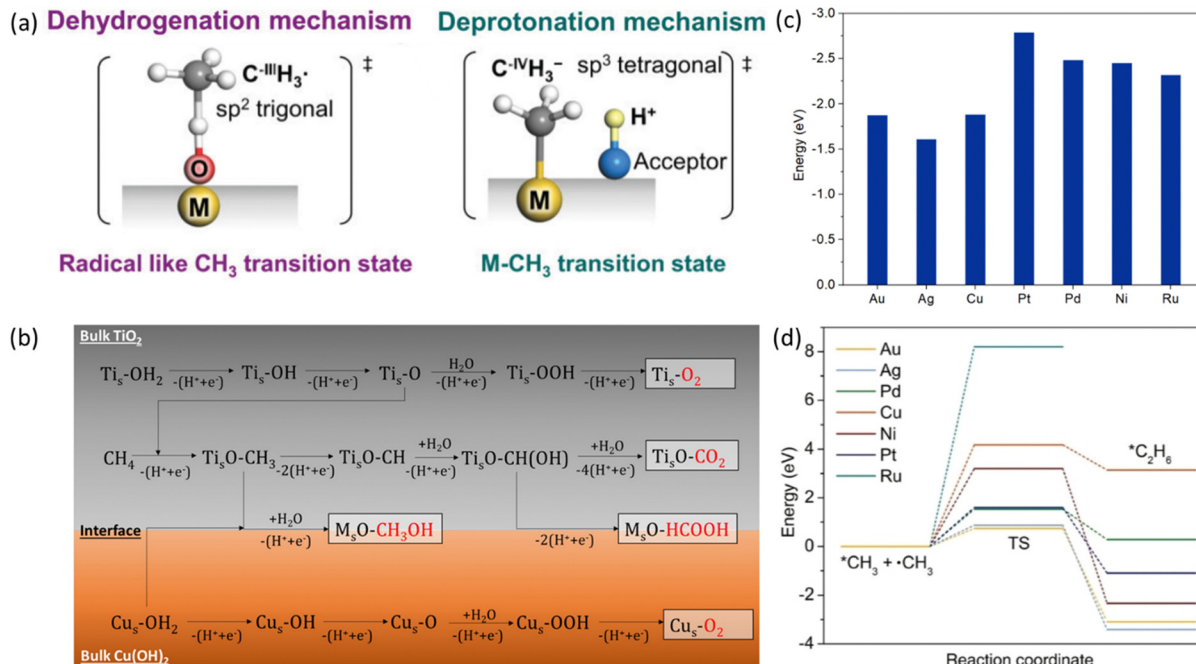


Fig. 3 (a) Dehydrogenation and deprotonation mechanisms in first C–H bond dissociation. Reproduced with permission.²⁶ Copyright 2020, Wiley–VCH. (b) Control of sequential C–H cleavage of the CH₄ molecule at the interface between TiO₂ and Cu(OH)₂. Reproduced with permission.²⁷ Copyright 2022 American Chemical Society. (c) Adsorption energy of *CH₃ on the surface of different metals. (d) Transition state energies for *CH₃ coupling reaction on the surface of different metals. Reproduced with permission.²⁸ Copyright 2023, Wiley–VCH.

in both EMC and PMC. To prevent this sequential dissociation, the *CH₃ groups need to be removed from the catalytic surfaces. The addition of cocatalysts can provide an interface that inhibits further dissociation. For example, CH₄ suffers from severe over-oxidation to CO₂ on a bare TiO₂ surface. With the addition of Cu(OH)₂ as a cocatalyst, it can provide additional reactive oxygen species (ROS) which can bind with *CH₃ to generate methanol preventing further C–H cleavage (Fig. 3b).²⁷ Besides ROS, *CH₃ can also go through a coupling process to form higher hydrocarbons on the metallic cocatalysts with a stronger adsorption energy of *CH₃ than it on metal oxides (Fig. 3c). Regarding C–C coupling, different metals exhibit varying behaviours due to differences in transition state energies, as shown in Fig. 3d.²⁸ These variations can serve as a selection criterion for the design of efficient catalysts for CH₄ coupling.

To understand the catalytic mechanism, different *in situ* characterization methods, *e.g.*, Raman spectroscopy, infrared Microscopy (IR), X-ray absorption spectroscopy (XAS), and electron paramagnetic resonance (EPR), can be employed together with theoretical calculations to confirm the reaction route. In the following sections, we will introduce how these different mechanisms can be applied in EMC and PMC investigations.

3 Electrocatalytic methane conversion

3.1 Mechanisms and recent advances

EMC represents a process to overcome the methane activation energy with external electric potential.³⁵ Depending on the

activation or reaction sites, methane activation can occur either on the electrode surfaces (*i.e.*, direct methane activation, Fig. 4a) or within the bulk liquid (*i.e.*, indirect activation, Fig. 4b). For indirect activation, it involves the generation of highly oxidative species in the bulk electrolyte, which will be consumed in the following methane oxidation process.¹⁵ The applied potential plays a key role in controlling the reaction rate. As shown in Fig. 4c, under applied potential (noted with overpotential, η), the methane adsorbed on electrodes can be significantly activated, leading to a much lower energy barrier ($\Delta G_F^\circ - \alpha z F \eta$) compared to the conventional thermocatalytic process. Meanwhile, the applied potential will also significantly reduce the free energy of the products within the reaction environment, as a result of which the EMC will encounter a much stronger driving force under applied bias.¹⁵ Moreover, the applied potential can be precisely controlled, leading to the adjustment of both activity and selectivity of the EMC process.¹⁹ All these features make EMC a promising approach for methane conversion.

In a typical EMC process under mild conditions (<50 °C), oxygenated species such as C₁–C₃ alcohols and carboxylic acids are the main products. During EMC, the active oxygen species surface coverage is significant for methane activation and substantially affect the final product selectivity and conversion rate.^{36–38} In heterogeneous EMC, water from the liquid electrolyte is the source for active oxygen species generation towards methane conversion products. Meanwhile, it will compete with the water oxidation for the oxygen evolution reaction (OER). Therefore, many strategies have been applied to facilitate EMC



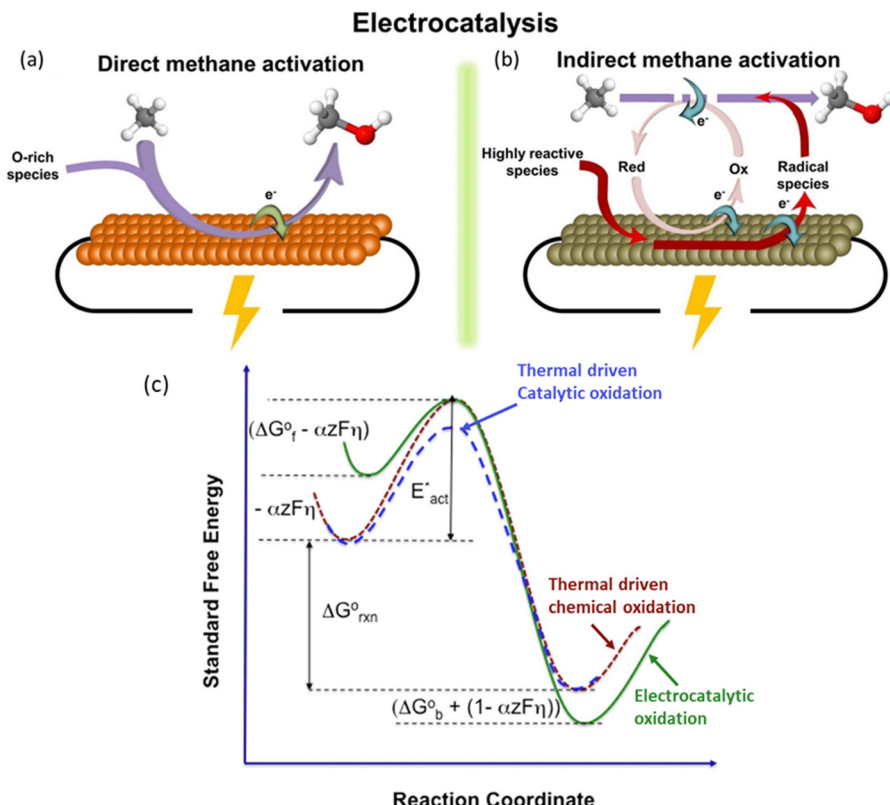


Fig. 4 Schematic illustration of (a) direct electrocatalytic methane activation, and (b) indirect electrocatalytic activation. Reproduced with permission.¹⁵ Copyright 2021, Elsevier. (c) Effects of overpotential on reaction energy profile. Reproduced with permission.¹⁹ Copyright 2016, Elsevier.

while suppressing the OER in electrocatalysis by materials engineering, electrolyte engineering, and/or more easily, precise control of applied bias. By contrast, homogeneous EMC provided an ultimate solution to suppress the OER by utilizing concentrated acidic electrolyte, which avoids aqueous content. These strategies are incorporated into recent advances of EMC, a summary of which is presented based on heterogeneous and homogeneous systems in Table 1.

3.2 Heterogeneous catalysts

Heterogeneous EMC originates from solid oxide fuel cell research using methane as a fuel. Generally, a methane conversion catalyst is directly deposited on a solid oxide electrolyte (SOE) as the ion conductor and thus results in an effective methane conversion. It can provide fast mass transfer of methane onto the catalytic surfaces with a gas diffusion layer. However, the activation of the SOE usually requires high temperature (over 200 °C), which is a bottleneck for the wide application of this method. Many efforts have been evoked to break this limitation with the innovation in SOE and development in efficient electrocatalysts, which meanwhile improved the selectivity towards specific product of interest.^{54–59} The very first EMC at nearly ambient temperature was performed on NiO/ZrO₂ with the use of a carbonate anion exchange membrane, where the interaction between ZrO₂ and CO₃²⁻ provides the key oxygen species for the reaction.³⁹ To make the reaction condition milder, studies are aimed at achieving ECM at room temperature with

a liquid electrolyte. ZrO₂ and CO₃²⁻ have been employed in combination to provide active oxygen species facilitating EMC and it has been reported that ZrO₂ anchored on oval-shaped cobalt(II, III) oxide (Co₃O₄) heterojunctions has supreme ability to react with CH₄.⁴⁰ Three-carbon (C₃) species products are generated following a proposed mechanism in Fig. 5a. The methane C–H bond dissociation undergoes a dehydrogenation mechanism, which is either followed by nucleophilic addition to form 2-propanol or free radical addition to form 1-propanol.⁴⁰ The following study uses bimetallic oxide NiCo₂O₄ to replace Co₃O₄ in a nanowire structure with similar heterojunctions between ZrO₂ and reaches a 65% product selectivity towards propionic acid.⁴¹ By further applying morphology control over the ZrO₂ substrate by forming a nanotube structure (Fig. 5b), the reaction rate is improved by 10-fold.^{40,42} Furthermore, CeO₂ shows low oxygen vacancy formation energy, leading to the enhanced active oxygen species generation from CO₃²⁻. Therefore, CeO₂-based materials were developed with monoclinic copper oxide (CuO), and the developed CeO₂/CuO has achieved significant methanol production with high selectivity.^{43,60}

EMC can be promoted by surface active oxygen species from CO₃²⁻ ions with the combination of material and electrolyte engineering. It is reported that the utilization of CO₃²⁻ ions can circumvent the complete OER and maintain a high surface coverage of active oxygen.²² In addition to carbonate, alkaline electrolytes including sodium hydroxide (NaOH) and potassium hydroxide (KOH) are also employed in EMC systems



Table 1 EMC under mild conditions

System	Catalyst	Reaction conditions	Electrolyte	Performance	Ref.
Heterogeneous	NiO/ZrO ₂	40 °C 2.0 V	1.0 M Na ₂ CO ₃ + DMF soaked AM-PAD anion exchange membrane	$J = 21 \text{ mA cm}^{-2}$	39
	ZrO ₂ /Co ₃ O ₄	2.0 V vs. Pt	0.5 M Na ₂ CO ₃	$J < 10 \text{ mA cm}^{-2}$ 1-Propanol: 111.3 $\mu\text{mol g}_{\text{cat}}^{-1} \text{ h}^{-1}$ 2-Propanol: 109.6 $\mu\text{mol g}_{\text{cat}}^{-1} \text{ h}^{-1}$ Production efficiency 60%	40
	ZrO ₂ /NiCo ₂ O ₄	2.0 V vs. Pt	0.5 M Na ₂ CO ₃	Conversion efficiency: 47.5% Propionic acid: 1173 $\mu\text{mol g}_{\text{cat}}^{-1} \text{ h}^{-1}$ Propionic acid selectivity: 65%	41
	Co ₃ O ₄ /ZrO ₂ NT ^a	2.0 V vs. SHE ^b	0.5 M Na ₂ CO ₃	1-Propanol: 2681 $\mu\text{mol g}_{\text{cat}}^{-1} \text{ h}^{-1}$ 2-Propanol: 1395 $\mu\text{mol g}_{\text{cat}}^{-1} \text{ h}^{-1}$ Product selectivity: 91.98%	42
	CuO/CeO ₂	1.5 V, 10 bar	0.5 M Na ₂ CO ₃	Methanol: 752.9 $\mu\text{mol g}_{\text{cat}}^{-1} \text{ h}^{-1}$ Methanol selectivity: 79%	43
	Fe ₃ Ni ₇ (OH) _x NS ^c	1.46 V vs. RHE ^d	0.1 M NaOH	Methanol: 1832.2 $\mu\text{mol g}_{\text{cat}}^{-1} \text{ h}^{-1}$ TOF = 936 Ethanol: 9090 $\mu\text{mol g}_{\text{cat}}^{-1} \text{ h}^{-1}$ faradaic efficiency: 87%	44
	NiO/Ni foam	1.4 V vs. RHE	0.1 M NaOH	Current efficiency = ~100% Ethanol selectivity: 89% Ethanol: 25 $\mu\text{mol g}_{\text{cat}}^{-1} \text{ h}^{-1}$ Methanol selectivity: 10% Methanol: 7.4 $\mu\text{mol g}_{\text{cat}}^{-1} \text{ h}^{-1}$ Ethanol batch: 4668.3 $\mu\text{mol g}_{\text{cat}}^{-1} \text{ h}^{-1}$ Ethanol flow: 11480.6 $\mu\text{mol g}_{\text{cat}}^{-1} \text{ h}^{-1}$ Ethanol selectivity: 85% Methanol selectivity: 15%	45
	Fe–N–C SAC ^e	1.6 V vs. RHE	0.1 M KOH	Production rate: 789 $\mu\text{mol g}_{\text{cat}}^{-1} \text{ h}^{-1}$ Ethanol selectivity: 85%	37
	Rh/ZnO NS	2.2 V vs. RHE	0.1 M KOH	Production rate: 127 $\mu\text{mol g}_{\text{cat}}^{-1} \text{ h}^{-1}$ Total FE: 10.9%	46
	Mg-MOF-74 ^f	1.6 V vs. RHE	1 M KOH	Production rate: 1680 $\mu\text{mol g}_{\text{cat}}^{-1} \text{ h}^{-1}$ Formic acid selectivity: 80.7%	47
Homogeneous	a-KB ^g	0 V vs. RHE	0.05 M H ₂ SO ₄	Production rate: 11.5 mmol g _{Fe} ⁻¹ h ⁻¹ TOF ^j = 2000	48
	Ag, Fe ^{II} _{III,III} Pd ₂	0.12 V vs. RHE 140 °C 34 bar	0.1 M HClO ₄ 20% oleum ⁱ	TOF = 2000	49
	Rh ^{III}	2.0 V vs. SSE ^h −1.4 V vs. SCE ^k	0.1 M TBACLO ₄ in 1,2 DFB	TOF = 2159	50
	V ₂ –oxo	2.255 V vs. SCE 3 bar	98% H ₂ SO ₄	TOF = 483 TOF = 1336	51
	Ag ^{II}	6 bar 1.737 V vs. MSE ^l	98% H ₂ SO ₄	TOF = 2800	52

If not stated, operation conditions are under ambient conditions of 25 °C, 1 bar. ^a NT: nanotube. ^b SHE: standard hydrogen electrode.

^c NS: nanosheet. ^d RHE: reversible hydrogen electrode. ^e SAC: single atom catalyst. ^f MOF: metal–organic framework. ^g a-KB: acid-treated Ketjen Black. ^h SSE: silver sulphate electrode. ⁱ 20% oleum contains 20% SO₃ and 80% H₂SO₄ by weight. ^j TOF: turnover frequency. ^k SCE: saturated calomel electrode. ^l MSE: mercurous sulphate electrode.

together with accurate control of applied bias to generate active oxygen species. The potential-dependent reaction mechanism usually leaves a small potential window that can simultaneously generate and stably maintain active oxygen species on the catalytic surfaces. In addition, methane overoxidation into CO₂ can also be prevented by adjusting the bias. This naturally makes accurate control of the applied bias on these catalytic surfaces crucial for achieving a high conversion of methane.

Taking the surface of a CoO_x film as an example, the EMC process consists of 4 major steps, including oxyhydroxide (−OOH) formation, methanol production, overoxidation to carbon dioxide, and the OER, respectively.³⁸ As shown in Fig. 6a, at potentials between 0.5 and 1.0 V vs. SHE, methane conversion

into methanol is dominant, while above 1.0 V vs. SHE, the overoxidation of methane into carbon dioxide is preferred, whereas the OER, which competes with methane in reacting with OH*, occurs and dominates at potential greater than 1.28 V vs. SHE.³⁸ Inspired by the above potential dependent reaction mechanism, iron–nickel hydroxide nanosheets (Fe₃Ni₇(OH)_x) are developed, which can convert methane into ethanol at 1.46 V vs. RHE, where the nickel oxyhydroxide (NiOOH) generated from NiO is considered as the active phase for the EMC process as revealed by the increased oxidation peak of Ni(II) to Ni(III) transition.^{44,45}

Accurate control of the applied bias on catalytic surfaces is crucial for achieving a high coverage of active oxygen species, given the potential-dependent OER process. Material engineering



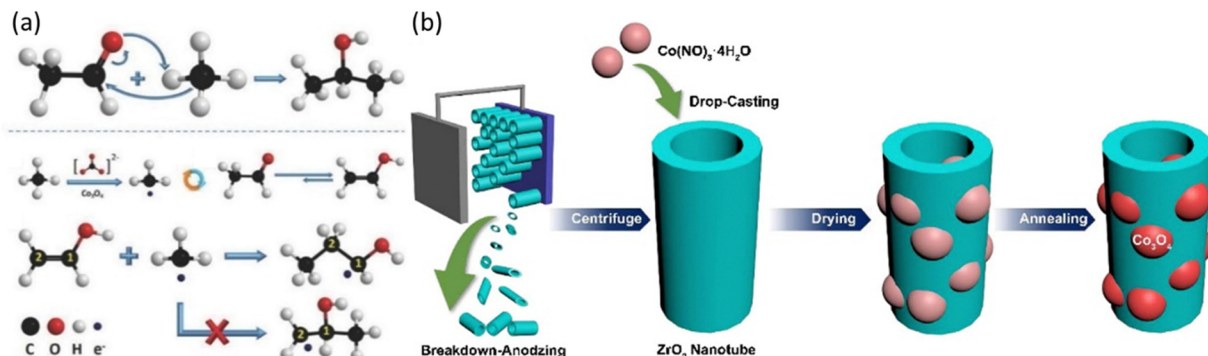


Fig. 5 (a) Nucleophilic addition reaction of methane to form 2-propanol and free radical addition reaction of methane with acetaldehyde to form 1-propanol. Reproduced with permission.⁴⁰ Copyright 2017, Wiley-VCH. (b) Schematic illustration of the ZrO_2 NT/ Co_3O_4 synthesis procedure. Reproduced with permission.⁴² Copyright 2021, Elsevier.

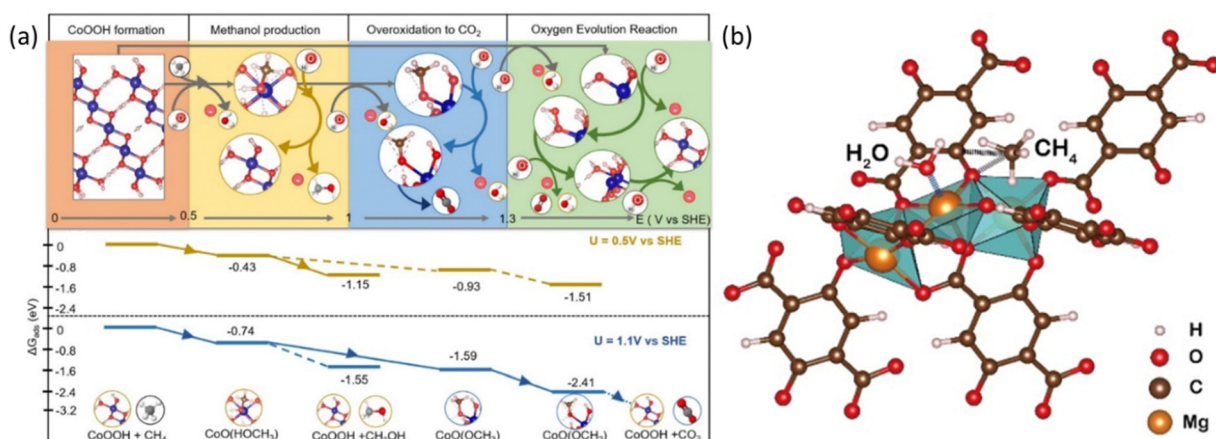


Fig. 6 (a) The top panel shows the favoured reactions at certain electrochemical potential range evaluated at $\text{pH} = 12$. The bottom panel shows reaction energy diagram at $U = 0.5 \text{ V}$ vs. SHE and 1.1 V vs. SHE. Reproduced with permission.³⁸ Copyright 2023, American Chemical Society. (b) Graphical illustration of the CH_4 adsorption on Mg-oxo-Mg and H_2O adsorption on Mg-MOF-74. Reproduced with permission.⁴⁷ Copyright 2022, Elsevier.

is employed to achieve a broad potential range that maximizes the generation of surface oxygen species without complete water oxidation *via* the OER pathway. For instance, on a Fe-N-C single-atom catalyst, active atomic oxygen can be stabilized even at relatively high potentials owing to its unique rate-limiting step in the OER.³⁷ Following a similar principle, rhodium-doped zinc oxide nanosheets (Rh/ZnO NS) and magnesium-substituted metal organic frameworks (Mg-MOF-74) have also demonstrated promising selectivity for EMC.^{46,47} Another strategy to avoid complete oxidation of water is to perform EMC with alternative oxidative agent rather than surface active oxygen species. For instance, EMC can be coupled with the oxygen reduction reaction (ORR) on cathodes. Such a strategy is successful on acid-treated ketjen black (a-KB) carbon powder working as the cathode catalyst for *in situ* ORR generating hydrogen peroxide (H_2O_2) (Fig. 7a), which subsequently generates active oxygen species to facilitate the EMC process.⁴⁸ The Fenton reaction for fast generating active oxygen species is then incorporated into such a system (Fig. 7b) with a silver foil cathode.⁴⁹ For this ORR-induced EMC process, the cathode is not directly

responsible for methane oxidation but generates active oxidative species reacting with methane molecules.

3.3 Homogeneous electrocatalysts

In homogeneous EMC systems, the CH_4 molecule is directly oxidized by the oxidative reagents produced on anodes.³² In these systems, highly oxidative intermediates will first be produced on anodes (*e.g.*, M^{n+2} in Fig. 8a), then the M^{n+2} will oxidize methane molecules into methanol (Fig. 8a). In the presence of concentrated H_2SO_4 , the electrophilic high valent metal ions can mediate methane conversion through a two-electron oxidation process into methyl esters, which can be protected from further being oxidized by the electron-withdrawing effect of the HSO_4^- groups. During these reactions, the capability for methane oxidation can be estimated by the redox potential of the metal ions (Fig. 8b).^{30,50} This strategy is usually performed in a non-aqueous electrolyte which wipes out the concern brought by the OER. Thus, the key research challenges in homogeneous EMC lie in the proper selection of



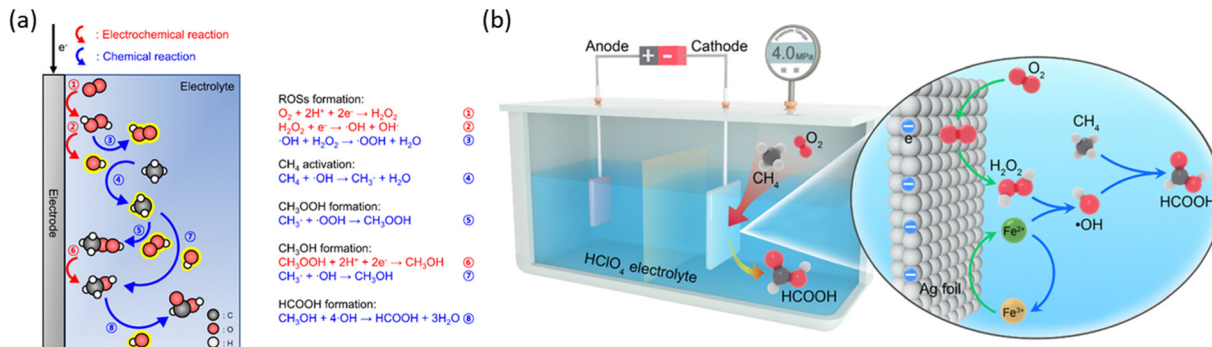


Fig. 7 (a) Schematic illustration of the reaction mechanism for direct POM with the *in situ* ORR. Reproduced with permission.⁴⁸ Copyright 2023, Springer Nature. (b) Schematic illustration of Fenton reaction assisted POM with the *in situ* ORR. Reproduced with permission.⁴⁹ Copyright 2024, American Chemical Society.

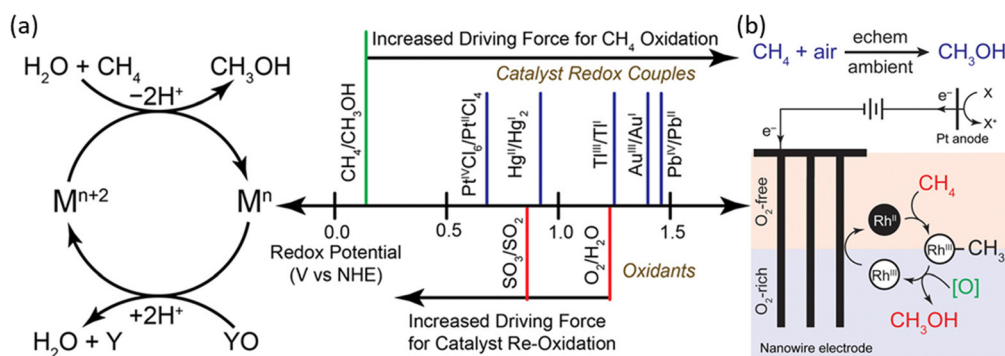


Fig. 8 (a) Redox potentials of methane oxidation catalysts. (Left) Simplified catalytic cycle for electrophilic methane oxidation using a stoichiometric oxidant such as SO₃ or O₂. (Right) Estimated redox potentials of electrophilic methane functionalization catalysts/reagents. Reproduced with permission.⁵⁰ Copyright 2017, American Chemical Society. (b) The proposed catalytic cycle of Rh^{II} functioning methane into methanol. Reproduced with permission.⁵¹ Copyright 2019, American Chemical Society.

redox couples for efficient methane conversion and stable redox loops.

Redox couples including Pd^{III,IV}, Ag^{I,II}, V₂-oxo, and Rh^{II,III} have been demonstrated to be efficient as homogeneous electrocatalysts (Table 1), in which Ag^{I,II} and V₂-oxo show the benchmark high CH₄ conversion rate. The V₂-oxo dimer is synthesised by dissolving V₂O₅ in 98% H₂SO₄ and can achieve EMC at a TOF of 1336 h⁻¹.⁵² The high efficiency of this catalyst is attributed to its extremely low activation energy of 10.8 ± 0.6 kcal mol⁻¹ in methane functionalization.⁵² The rate-limiting step is then considered either the dissolution of methane or the one-electron oxidation of the V-oxo dimer.⁵² Another work on Ag^{I,II} achieved a TOF of 2800 h⁻¹ in 3 h at ambient temperature. Moreover, one unique approach reported is by utilizing Rh^{II} tetramesitylporphyrin metalloradicals, which functionalised CH₄ via the oxidation of Rh^{II} to Rh^{III}, as illustrated in Fig. 8b.

The EMC driven by a homogeneous process with different redox couples can avoid the OER process and achieve a fast CH₄ oxidation. However, the utilized highly acidic electrolyte presents as a high-risk, which also makes the long-term stability a concern during operation. The efforts in homogenous EMC should be mainly devoted to lower the harshness of the

reaction conditions. Despite these advances, EMC is hindered by its low efficiency because of the low solubility and widespread product distribution. Future development in this field necessitates a focus on the development of novel catalyst design, such as bifunctional electrocatalysts, to provide active sites for reactant adsorption and the dissociation of methane C-H bonds.

4 Photocatalytic methane coupling into ethylene

The electrocatalytic methane conversion generally requires a liquid phase electrolyte, which will lead to a significant mass diffusion issue at high current application. To overcome this issue, gaseous phase CH₄ conversion is more applicable for high-rate methane conversion. Photocatalytic methane conversion (PMC) is considered the most cutting-edge strategy for high efficiency and green methane conversion. In the research community, some remarkable advances have been made in methane coupling to ethane. However, for the more promising product, ethylene, there has been no good summary in revisiting the reaction mechanism and photocatalyst design. In the

following section, we will focus on the recent research progress on PMC for ethylene generation.

4.1 Mechanisms and pathways

PMC offers a promising solution by harvesting the energy from sunlight using semiconductor materials.⁶¹ As shown in Fig. 9, under light irradiation with energy greater than the energy gap of the semiconductor, the electrons in the valence band (VB) will be excited to the conduction band (CB) leaving vacancies regarded as holes in the VB. The photogenerated electrons and holes will then transfer to the semiconductor surface and facilitate the redox reactions. To drive CH_4 conversion, the position of the VB in the semiconductor should be more positive than the redox potential required for breaking C–H bonds (+2.06 V vs. SHE); however the position of the CB should be more negative than the redox potential required for reducing O_2 in the OCM or the redox potential for H_2 production for NOCM (0 V vs. SHE).^{23,62} Due to the abundance and eco-friendliness of photoenergy on Earth, this promising solution not only provides the potential to revolutionize ethylene production but also solves the increasing demand for cleaner and cost-effective methods in the industry.⁶³

During PMC, it is regarded that CH_4 is first oxidized by photogenerated holes to form a methyl intermediate ($^*\text{CH}_3$) which then can be further coupled together to form C_2 products (e.g., C_2H_6 and C_2H_4).¹¹ Simultaneously, in photocatalytic NOCM (Fig. 9a), protons (H^+) are formed and further reduced by photogenerated electrons producing hydrogen, whereas in photocatalytic OCM (Fig. 9a), O_2 is reduced by photogenerated electrons together with protons to form water. Ethylene production can be achieved through two different possible pathways followed by the first C–H bond dissociation of methane as shown in Fig. 10.⁶⁴ One pathway is that the $^*\text{CH}_3$ will first couple together to form C_2H_6 and then produce C_2H_4 via a dehydrogenation process. Alternatively, the formed $^*\text{CH}_3$ will be further cleaved to form methylene intermediate ($^*\text{CH}_2$) for the coupling reaction for C_2H_4 generation. Despite the above assumptions, currently, there is still no consensus on the most plausible pathway for C_2H_4 generation. This reaction can be affected by multiple factors, such as photocatalyst design, cocatalyst loading, different oxidative agents for OCM, and reactor configuration. The following sections discuss how these factors affect the C_2H_4 production *via* PMC.

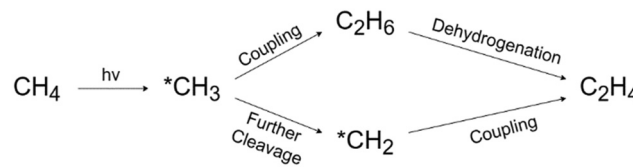


Fig. 10 Possible reaction pathways for ethylene production.

4.2 Semiconductor photocatalyst design

Semiconductors are essential components for absorbing light and producing photogenerated electron–hole pairs during PMC. Metal oxide semiconductors, such as TiO_2 , WO_3 , and ZnO , are widely used in photocatalytic methane conversion, due to the much positive position of O 2p orbitals (approximately at +3 V vs. SHE), which has a strong oxidation capability upon light irradiation to produce reactive oxygen species (ROS) (e.g., O^- and O_2^-).⁶⁵ These ROSs are highly active in cleaving the protons from the C–H bond in CH_4 *via* the electrophilic activation process. However, it is noteworthy mentioning that under a gaseous environment, the above redox potential may not be applicable, which is dedicated to the aqueous environment, and more investigations are required for clarifying the selection criteria for potential semiconductors. Table 2 summarizes the published works on photocatalytic coupling CH_4 to C_2H_4 .

In the early period, gallium oxide was researched as a photocatalyst for NOCM in which a little amount of C_2H_4 ($0.005 \mu\text{mol g}^{-1} \text{h}^{-1}$) was observed with a carbon selectivity of only 2.8%.⁶⁶ C_2H_4 was identified as an adsorbed product on the catalysts, which can only be obtained after desorbing at 300 °C. Similarly, using supported ceria photocatalysts (as shown in Fig. 11a with other lanthanoid elements), trace amounts of C_2H_4 production can be observed directly without the desorbing process, while the majority of the C_2H_4 product ($0.1 \mu\text{mol g}^{-1} \text{h}^{-1}$) with 22.7% selectivity was obtained by thermal desorption on the low ceria loading samples.⁶⁷ Attention should be paid to determine whether the source of the C_2H_4 product is from photocatalysis or thermal catalysis. It is claimed that the main species on the low-loading sample were $\text{Ce}(\text{III})$ oxide isolated monomers with high dispersity on the surface of the support (*i.e.*, SiO_2 and Al_2O_3) due to the quantum size effect. According to the photocatalytic performance and its methane quenching experiments, these highly dispersed $\text{Ce}(\text{III})$ species would act as the active sites for photocatalytic NOCM. Nevertheless, pure intrinsic semiconductors

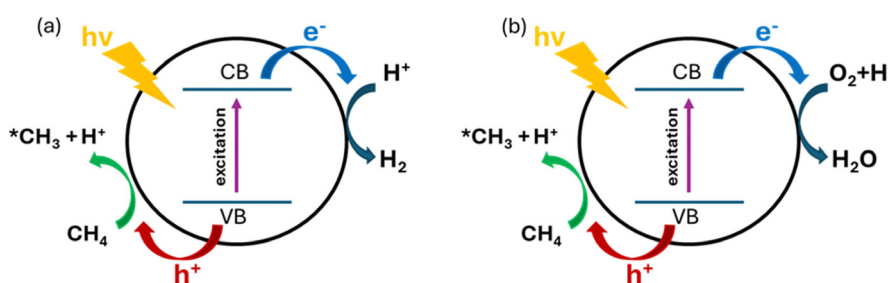


Fig. 9 Schematic representation of photocatalytic (a) NOCM, and (b) OCM.



Table 2 Photocatalytic coupling of CH_4 to C_2H_4

Photocatalyst	Cocatalyst	C_2H_4 production rate ($\mu\text{mol g}^{-1} \text{h}^{-1}$)	C_2H_4 selectivity	Reaction conditions	Ref.
$\text{Ga}_2\text{O}_3\text{-P}$	—	0.005	2.8%	300 W Xe lamp; 310 K; 3 h; 0.2 g catalyst; 200 μmol CH_4	66
Ce/SiO_2	—	0.05	14.6%	300 W Xe lamp; 310 K; 3 h; 0.2 g catalyst; 200 μmol CH_4	67
$\text{Ce/Al}_2\text{O}_3$	—	0.1	22.7%	300 W Xe lamp; 310 K; 3 h; 0.2 g catalyst; 200 μmol CH_4	67
Ga ³⁺ -modified Titanosilicate (ETS-10)	—	0.5	3.8%	150 W high-pressure Hg lamp; 5 h; 0.2 g catalyst; 200 μmol CH_4	68
TiO ₂ nanotubes	—	54.58	50.2%	300 W Xe lamp; 4 h; 3 mg catalyst; 5 mL CH_4	69
ZnO	Au	Trace amount	N/A	365 nm LED (PLS-LED100C); 4 h; 5 mg catalyst; $\text{CH}_4/\text{O}_2 = 99/1$, 56 mL gas	28
In ₂ O ₃	Ag	Trace amount	N/A	300 W Xe lamp; 1 h; 20 mg catalyst; 175 mL CH_4	70
TiO ₂ (P25)	Pt	1.11	2%	UV lamp (Philips, TUV 4W/G4 T5; wavelength at 254 nm); 6 h; 75 mg catalyst; 80 mL CH_4	71
TiO ₂ (PC-50)	Pt, CuO_x	2.4	2.1%	40 W 365 nm LED; 100 mg catalyst; $\text{O}_2:\text{CH}_4 = 1:400$; GHSV = 2400 h^{-1} ; 10% CH_4 ; Flow reactor	72
ZnO	$\text{AuPd}_{2.7}$	13.3	38.1%	300 W Xe lamp; 8 h; 2 mg catalyst; 0.5 mL CH_4	73
WO ₃	$\text{Pd}_5/\text{Zn}_{0.35}$	24	75.3%	300 W Xe lamp; 2 h; 2 mg catalyst; 0.5 mL CH_4	74
ZnO/TiO ₂	Au	20	0.36%	300 W Xe lamp; 20 mg catalyst; $\text{CH}_4/\text{synthetic air}$ (20 vol% O_2/N_2) = 69/1; Flowrate = 70 mL min^{-1} ; Flow reactor	75
Bi ₂ Nb ₅ F	$\text{Au}_2\text{-Pd}_2$	22.6	63%	300 W Xe lamp; 2 h; 5 mg catalyst; 1 mL CH_4	76
TiO ₂ (P25)	Ag	686	54.4%	Xe lamp (Power of 84.2 mW cm^{-2}); 2 h; 100 mg catalyst; $\text{CO}_2/\text{CH}_4/\text{Ar} = 7.5/7.5/85$; 2 MPa	77
TiO ₂ (PC-50)	PdCu Nanoalloy	60	1.25%	40 W LED 365 nm; 50 mg catalyst; $\text{CH}_4/\text{O}_2 = 114/1$, 10% CH_4 (Ar in Balance); GHSV = 342 000 mL g^{-1} hour ⁻¹ ; Flow reactor	78
TiO ₂	Sputtered Au	140	0.5%	100 W LED 365 nm; 20 mg catalyst; 320 mL min^{-1} CH_4 , 12 mL min^{-1} air; 393 K; Flow reactor	79
Carbon-doped ZnO	Au	45.85	90.79%	300 W Xe lamp; 3 h; 100 mg catalyst; 1.5% CH_4 (Ar in balance), 500 mL gas	80
ZnO	Ag	19.74	10.53%	300 W Xe lamp; 0.5 g catalyst; 5% CH_4 (N_2 in balance); Flowrate = 10 mL min^{-1} ; Flow reactor	81
TiO ₂	Au0.05-Pd0.05	794	12.11%	300 W Xe lamp; 10 mg catalyst; moist CH_4 70 mL min^{-1} ; Flow reactor	82
TiO ₂	Pd	22	2.4%	300 W Xe lamp; 3 h; 3 mg catalyst; 30 mL CH_4	83

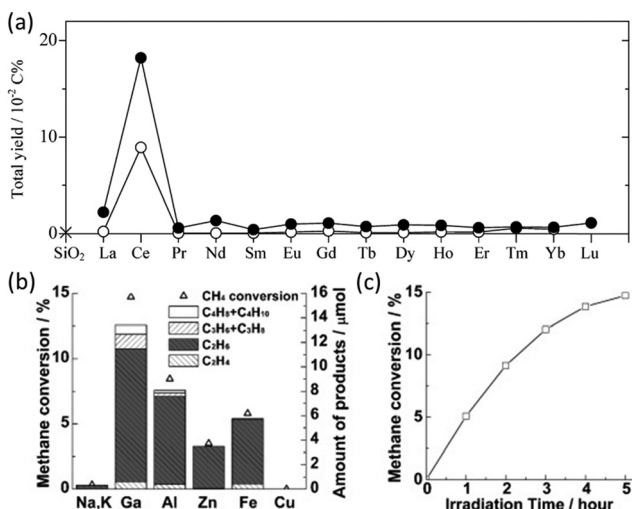


Fig. 11 (a) Total hydrocarbon yield in photocatalytic NOCM over SiO_2 (cross), lanthanoid elements (0.1 mol%) decorated SiO_2 (open circles), and lanthanoid elements (2 mol%) decorated SiO_2 (closed circles); Reproduced with permission.⁶⁷ Copyright 2008, American Chemical Society; (b) Photocatalytic NOCM performance over ETS-10 doped with different metal ions; (c) Photocatalytic NOCM performance as a function of time over Ga-ETS-10. Reproduced with permission.⁶⁸ Copyright 2012, Wiley-VCH.

suffer from rapid recombination of photogenerated electron–hole pairs and insufficient active sites.

To avoid the charge recombination possibility and enhance the activation of C–H bonds, a doping strategy has been incorporated. A study focusing on titanosilicate (ETS-10) with a microporous structure containing titanate semiconductors surrounded by the SiO_2 matrix used different metal ion dopants to enhance its photoactivity.⁶⁸ Compared with the intrinsic EST-10, the metal-modified (*i.e.*, Ga, Al, Zn, and Fe) EST-10 exhibited a dramatic enhancement on the photocatalytic NOCM performance due to the interaction of methane and the binary active species (*i.e.*, dopant metal ions and photo-generated holes).⁸⁴ Among them, Ga-modified EST-10 showed the best NOCM performance with almost 15% methane conversion rate (Fig. 11b and c) due to the strong C–H polarization ability.

In addition to metal doping, non-metal doping has been researched recently. Carbon-doped ZnO was constructed for efficient photocatalytic NOCM to C_2H_4 at a rate of 45.85 $\mu\text{mol g}^{-1} \text{h}^{-1}$ with excellent selectivity of 90% for C_2H_4 production and stoichiometric H_2 (88.07 $\mu\text{mol g}^{-1} \text{h}^{-1}$). The carbon doping in ZnO was proven to improve the stability of lattice oxygen, light absorption, and CH_4 activation.⁸⁵ This photocatalyst can not only weaken excitonic confinement to enhance charge separation but also suppress the overoxidation of hydrocarbons.⁸⁶ To achieve efficient C_2H_4 production, this photocatalyst generates $\text{Zn}^{+}\text{O}^{-}$ pairs to cleave C–H bonds, stabilizes the methoxy intermediate ($^{*}\text{OCH}_3$) for C–C coupling, and promotes the low-valence Zn generation for



dehydrogenation of the formed ethoxy intermediate ($^*\text{OC}_2\text{H}_5$) to produce C_2H_4 .

Crystal facet engineering has also been proven to be a promising strategy to fine-tune the structure of the photocatalysts by exposing the highly active facets to optimize the performance.^{87–89} Furthermore, defect engineering provides an alternative way to boost the photocatalyst performance by extending the light absorption range, increasing the surface active sites, and stabilizing key intermediates for C–H cleavage.⁹⁰ A recent study combined these two strategies (Fig. 12a) to construct oxygen vacancy self-doped single crystal-like TiO_2 nanotubes (Vo-p-TNT) with preferential crystalline orientations (001).⁶⁹ Through the modification of crystal facet and oxygen vacancy, this material achieved a C_2H_4 production rate of $54.58 \mu\text{mol g}^{-1} \text{h}^{-1}$ with 50.2% selectivity (Fig. 12b). Due to the coke deposition, the stability of this material was mediocre (Fig. 12c). The preferential crystal facet (001) was proved to improve the photogenerated charge separation and transfer, whereas the oxygen vacancy offers a significant number of unsaturated coordination sites for CH_4 adsorption and acts as electron traps to avoid charge recombination. The synergistic effect of oxygen vacancy and (001) facets resulted in an effective pathway for electron transfer between the photocatalyst and the adsorbed CH_4 molecule.⁹¹ The crystal facet engineering implemented on other photocatalysts (such as ZnO and WO_3) rather than TiO_2 requires exploration.

4.3 Cocatalyst loading

Loading cocatalysts is an efficient way to improve the photocatalytic performance.⁹² In a PMC system, the cocatalysts have several advantages, including (1) improving charge separation

and transfer efficiency; (2) providing active sites for CH_4 activation; (3) enhancing the stability of photocatalysts; (4) controlling the selectivity of desired products.⁹³ Currently, the research on cocatalysts in the field of PMC majorly focuses on noble metals, *e.g.*, Pt, Ag, Pd, and Au.²⁴

During PMC, CH_4 will firstly be oxidized to $^*\text{CH}_3$ by photo-generated charges. With the loading of metal cocatalysts, the formed $^*\text{CH}_3$ can be trapped on the metal surface from the photocatalyst and further couple together to produce C_2 products as shown in Fig. 13a. Various metallic cocatalysts (*e.g.*, Au, Ag, Pd, Cu, Ni, Ru, and Pt) loaded on the surface of the ZnO photocatalyst *via* the chemical reduction method have been studied for PMC (Fig. 13b).²⁸ Although the major product was C_2H_6 with C_2H_4 production as a minor reaction, the mechanistic study presented the uniqueness of Au in photocatalytic OCM compared with other metallic cocatalysts. The photogenerated $^*\text{CH}_3$ tended to be coupled on Au but over-oxidized on other metals due to the small coupling energy barriers (0.74 eV) on Au and the strong d–σ hybridization energy state (−5.63 eV) between Au and $^*\text{CH}_3$, which is below the Fermi level of Au.⁹⁴ Moreover, Au loaded on photocatalysts can promote the adsorption and activation of O_2 and increase the number of active photogenerated holes.

In addition to Au, other noble metals (*e.g.*, Pt, Pd, and Ag) have also been investigated in PMC. Ag offers the second smallest coupling energy barrier (0.88 eV). Moreover, nanosized Ag can provide the localized surface plasmon resonance (LSPR) on semiconductor photocatalysts inducing visible light response.⁹⁵ Due to this effect, TiO_2 -supported Ag nanoparticles were synthesized for the photocatalytic OCM reaction (Fig. 13c),

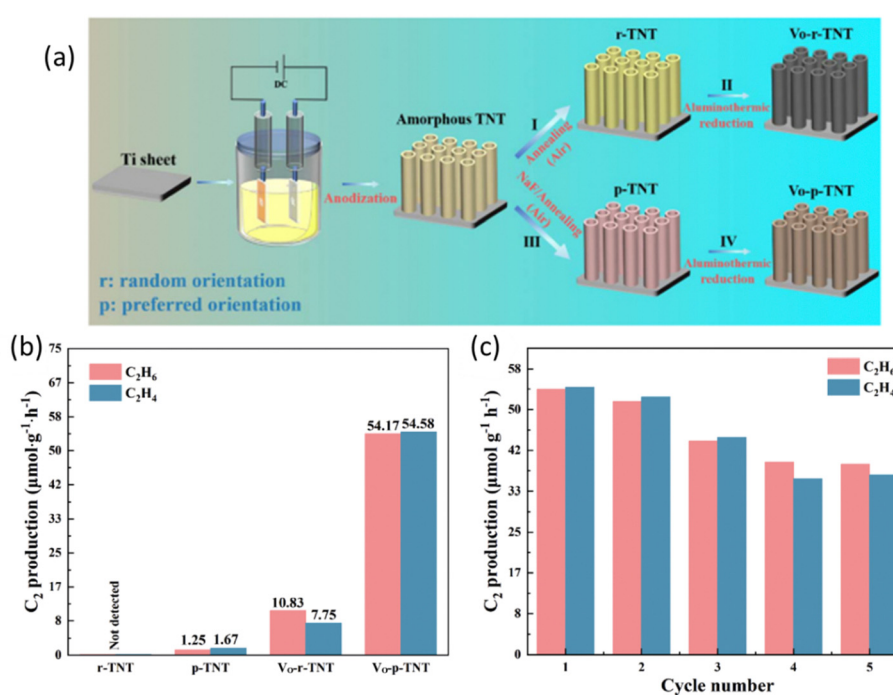


Fig. 12 (a) Schematic representation of the preparation process for different types of TNT; (b) photocatalytic NOCM performance over different types of TNT; (c) cyclic photocatalytic NOCM tests of Vo-p-TNT. Reproduced with permission.⁶⁹ Copyright 2023, Tsinghua University Press Ltd.



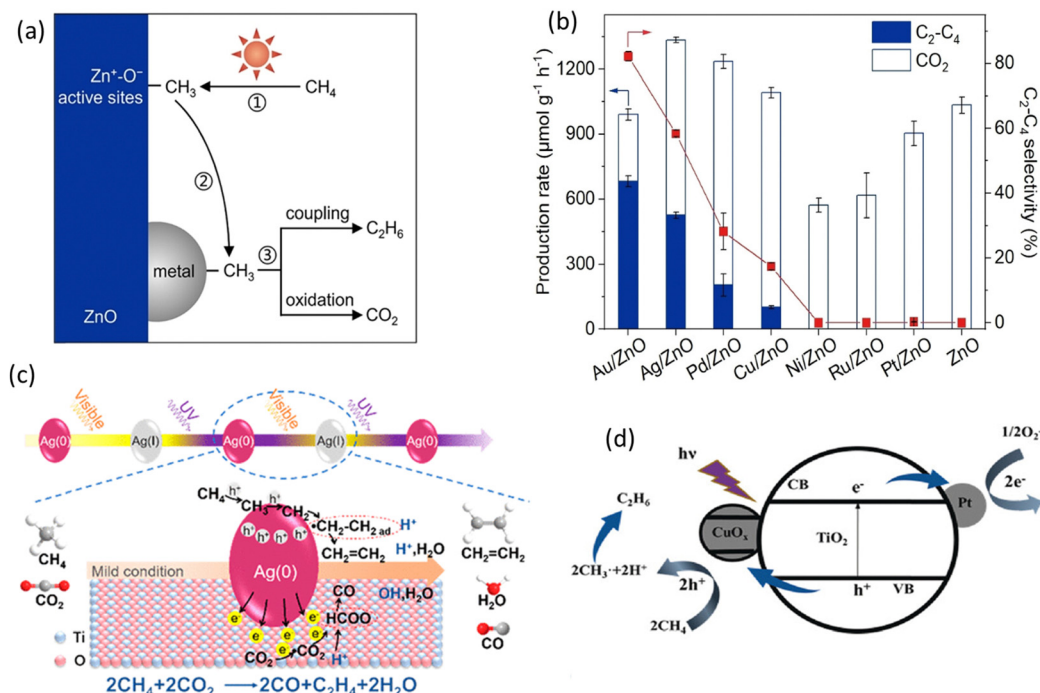


Fig. 13 (a) Schematic representation of the photocatalytic OCM reaction mechanism on metal/ZnO; (b) Photocatalytic OCM performance over ZnO and different metal loaded ZnO. Reproduced with permission.²⁸ Copyright 2023, Wiley-VCH. (c) Schematic representation of the photocatalytic CH₄ and CO₂ to C₂H₄ reaction mechanism over Ag/TiO₂; Reproduced with permission.⁷⁷ Copyright 2019, American Chemical Society. (d) Schematic representation of the photocatalytic OCM mechanism over Pt and CuO_x decorated TiO₂; Reproduced with permission.⁷² Copyright 2020, Wiley-VCH.

which showed a very high C₂H₄ production rate of 686 μmol g⁻¹ h⁻¹ with a selectivity of 54.4%.⁷⁷ The Ag nanoparticle was activated by visible light generating hot electrons and holes, which makes the CH₄ activation easier but leads to oxidation of Ag(0) to Ag(I) and then recovered by UV light. This work provided a feasible photocatalytic CH₄ to C₂H₄ production rate. Pt is a well-known electron acceptor when decorated on the surface of semiconductor photocatalysts.⁹⁶ Together with copper oxide (CuO_x), Pt was decorated on the surface of TiO₂ (Fig. 13d), showing a high yield for C₂ hydrocarbons with more than 60% selectivity in photocatalytic OCM in a flow reactor.⁷² With only Pt, the C₂ selectivity was decreased, while the CO₂ selectivity was increased. Pt as an electron acceptor facilitates charge separation, whereas CuO_x as a hole acceptor avoids overoxidation due to its less positive valence bands than TiO₂. The dual cocatalyst improved the C₂ yield by approximately 3.5 times higher than the bare TiO₂ photocatalyst. Similar to Pt, Pd nanoparticles also have a high oxidation ability.⁹⁷ A single atom cocatalyst was designed to avoid overoxidation in photocatalytic NOCM due to the stabilization of the lattice oxygen by Pd–O coordination altering the valence band maximum (VBM) of TiO₂.⁸³ Nevertheless, the product was majorly C₂H₆ rather than C₂H₄.

Compared to ethane, C₂H₄ is a more valued chemical but more challenging to be obtained in PMC. Inspired by thermocatalysis and electrocatalysis,^{98,99} Au and Pd nanoparticles were loaded on a BiNbO₅F photocatalyst to create a cascade reaction for C₂H₄ production *via* a two-step process, *i.e.*, activation of CH₄ on Au to produce C₂H₆ and dehydrogenation of C₂H₆ on Pd to C₂H₄, as shown in Fig. 14a.⁷⁶ This photocatalyst produced

C₂H₄ at a rate of 22.6 μmol g⁻¹ h⁻¹ with 63% selectivity. This work not only presents a way to produce C₂H₄ from CH₄ but also provides a general strategy for the cocatalyst design to efficiently and selectively drive complex reactions based on the tandem system. In addition to Au and Pd, Zn and Pd bimetallic cocatalysts over WO₃ photocatalysts were researched for photocatalytic NOCM to C₂H₄, showing a C₂H₄ production rate of 24 μmol g⁻¹ h⁻¹ with excellent selectivity (75.3%).⁷⁴ Based on the characterization results, the incorporation of Zn facilitated the cleavage of C–H bonds from CH₄ to form *CH₃ and the subsequent coupling, while Pd provided active sites to further dehydrogenate *CH₃ to *CH₂. The synergistic effect of Pd and Zn in WO₃ photocatalysts resulted in the highly selective C₂H₄ production from CH₄, inhibiting overoxidation to CO and CO₂.

Formation of metallic nanoalloy cocatalysts can also be used to boost C₂H₄ production. To overcome the weak dehydrogenation capability of Au, PdAu nanoalloys were synthesized on the surface of ZnO in which the highly dispersed Pd atoms within the Au lattice played a vital role, as shown in Fig. 14b. Au can facilitate photogenerated charge carrier separation, whereas Pd can induce further dehydrogenation capability.¹⁰⁰ The CH₄ molecules were first activated on the surface of the ZnO undergoing the dissociation process forming *CH₃ which would then react with lattice oxygen to form *OCH₃. With the help of Pd atoms, the *OCH₃ can be dehydrogenated to *CH₂O, which can then react with another CH₄ molecule to generate an ethoxy intermediate (*OC₂H₅). In the end, the *OC₂H₅ will be further dehydrogenated to produce C₂H₄ by Pd atoms. These findings

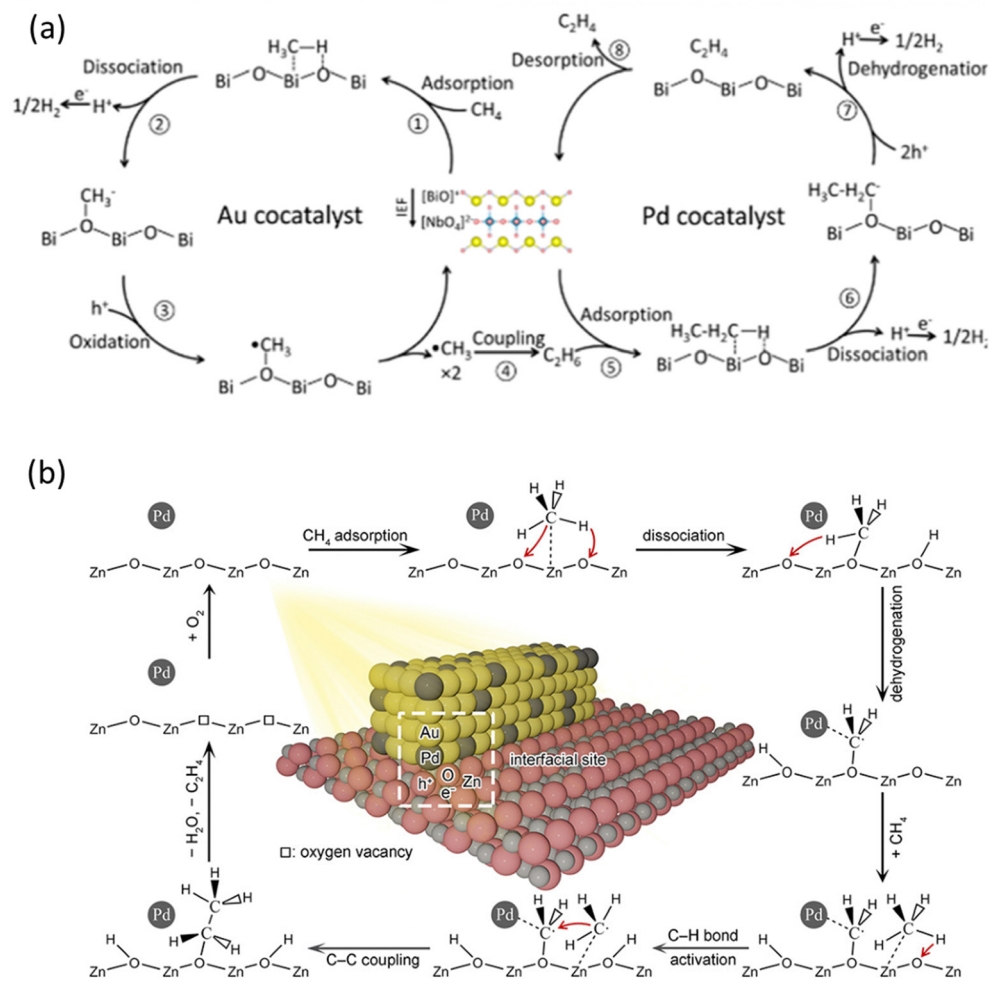


Fig. 14 Schematic representation of photocatalytic NOCM to C_2H_4 reaction mechanisms over (a) Au-Pd/BiNbO₅F; Reproduced with permission.⁷⁶ Copyright 2023, American Chemical Society. (b) AuPd/ZnO. Reproduced with permission.⁷³ Copyright 2020, American Chemical Society.

suggest the interaction between Pd atoms and $^{\bullet}OCH_3$ to promote the dehydrogenation process.

4.4 Reactor configuration

Rational reactor design is crucial to achieve efficient and selective PMC activity and selectivity.¹⁰¹ Batch and flow reactors are two main types of photoreactors, as shown in Fig. 15a and b, respectively. In the batch reaction system, methane and/or oxidants are introduced into a sealed reactor where the photocatalyst is present. As the reaction progresses, the products generated cannot be promptly removed, leading to overoxidation to CO_2 . Given that the free energy of the methane coupling products is generally lower than that of methane itself,¹⁰² achieving thermodynamically favourable conditions to desired products with high yield and selectivity presents a formidable challenge within the confines of a batch reactor.¹⁷ Furthermore, mass transfer limitations are observed in gas-solid reactions within batch reactors.¹⁰³ To overcome the limitations, two strategies have been introduced, *i.e.*, circulation process and increasing the pressure of the system. The circulation process utilizes a pump to improve the mass transfer and ensures a

homogeneously mixed gas environment. Increasing the pressure of the system can enhance the interaction between methane and the photocatalysts. According to Le Chatelier's principle, the reaction will shift towards decreasing the amount of gas (*i.e.*, CH_4 coupling) when the reactor pressure increases. For example, using CO_2 as a mild oxidant, Ag/TiO_2 achieved a high C_2H_4 production rate of $686 \mu\text{mol g}^{-1} \text{h}^{-1}$ under high pressure (*i.e.*, 2 MPa).⁷⁷ This indicates the significant impact of pressure increase.

On the other hand, flow reactors provide certain advantages over batch reactors as they improve mass transfer and enable adjustments in residence time and molecule–catalyst interactions, affording precise control over the interactions between methane and photocatalysts.⁷⁵ With the timely removal of the products, the overoxidation reaction is suppressed. The single-run conversion rate assessment stands out as a particularly valuable reference for further applications.¹⁷ Notably, a flow system provides a more uniform dispersion of methane and other reactant molecules than batch reactors.¹⁰⁴ Given these intrinsic advantages, assessing photocatalytic activity of methane coupling is critical. In an example of flow reactor utilization in



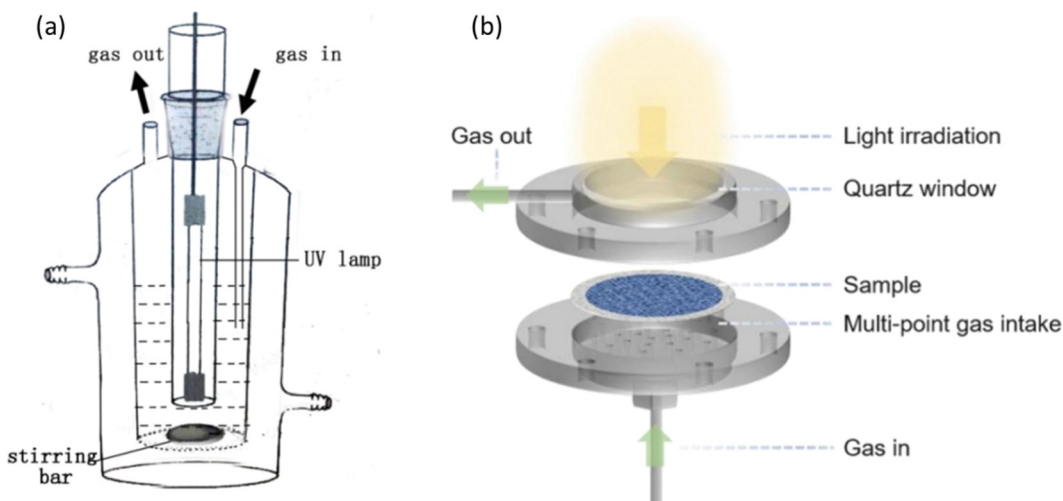


Fig. 15 Schematic representation of photocatalytic methane conversion in (a) batch reactor; reproduced with permission.⁷¹ Copyright 2016, Elsevier. (b) Flow reactor. Reproduced with permission.¹⁰⁴ Copyright 2024, American Chemical Society.

photocatalytic OCM reaction, the Au-ZnO/TiO₂ hybrid photocatalyst was employed.⁷⁵ Compared with its performance under batch reaction, the C₂ product selectivity was significantly enhanced under flow reaction. The rapid flow conditions lead to the migration of methyl radicals for coupling, which prevents photogenerated holes from oxidizing them further and reduces the yield of unwanted byproducts (e.g., coke, CO and CO₂).¹⁷ Compared to the batch reaction system, these combined results highlight the advantages of a flow reaction system.

4.5 Photocatalytic CH₄–C₂H₆–C₂H₄ system

Most of the aforementioned photocatalysts applied in photocatalytic methane coupling reactions present a pronounced tendency to yield C₂H₆ as the primary product, while the production of C₂H₄ plays a minor role. Attaining high selectivity towards C₂H₄ from photocatalytic CH₄ coupling is still a challenging endeavor. Presently, industrial synthesis of C₂H₄ involves steam cracking, utilizing C₂H₆ as the principal feedstock. Drawing inspiration from this conventional industrial process, a cascade photocatalytic system denoted as the photocatalytic CH₄–C₂H₆–C₂H₄ system is suggested to be implemented (Fig. 16). The photocatalytic dehydrogenation of C₂H₆ plays an important role in this system, which will be reviewed in this section. Upon light irradiation, the photogenerated holes in the semiconductor surface will attack the C–H bond in C₂H₆ molecules producing ethyl radicals (*C₂H₅) which then can be spontaneously dehydrogenated into C₂H₄ due to its intrinsic instability.^{105–107} Table 3 summarizes the works on photocatalytic dehydrogenation of C₂H₆ to C₂H₄.

The first work of photocatalytic C₂H₆ dehydrogenation to C₂H₄ employed a Pd-deposited TiO₂ (P25) photocatalyst using CO₂ as the mild oxidant (Fig. 17a).¹⁰⁸ This photocatalyst achieved a C₂H₄ production rate of 230.5 μmol g⁻¹ h⁻¹ with 95.42% selectivity. Pd was proved to lower the VB and CB of TiO₂, increasing the oxidation ability of the photogenerated holes upon light irradiation. The role of Pd was further

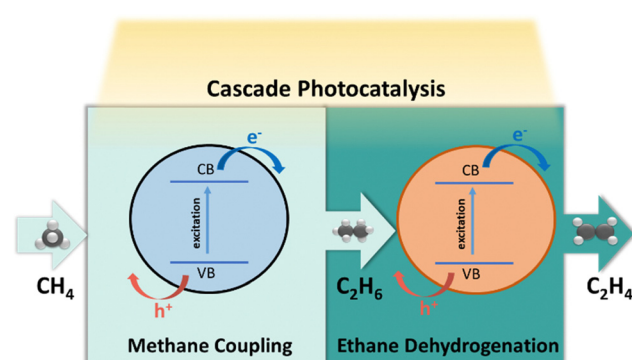


Fig. 16 Schematic representation of the photocatalytic CH₄–C₂H₆–C₂H₄ cascade system.

evidenced by another study investigating the effect of particle size.¹¹² Furthermore, PdZn intermetallic nanoparticles supported on ZnO with the robust interface to promote C₂H₆ activation were recently reported with an excellent C₂H₄ production rate of 46.4 mmol g⁻¹ h⁻¹ with 92.6% C₂H₄ selectivity using O₂ as the oxidant. These works demonstrated that photocatalytic oxidative dehydrogenation of C₂H₆ is a promising approach for C₂H₄ production from C₂H₆-containing feedstock.

Photocatalytic non-oxidative C₂H₆ dehydrogenation has also been researched using Pt-deposited LaVO₄ photocatalysts with rich oxygen vacancies through a dynamic synergistic effect of lattice oxygen and oxygen vacancies (Fig. 17b).¹¹⁰ This photocatalyst also presented excellent anti-coking ability with a C₂H₄ production rate of 275 μmol g⁻¹ h⁻¹ and 96.8% selectivity. The bimetallic system using Pd–Rh with internal electron transfer was also utilized to improve the light absorption and reactant adsorption.¹¹¹ Besides, a non-noble metal based photocatalyst using Cu-deposited TiO₂ (P25) was also used, producing C₂H₄ at a rate of 533.46 μmol g⁻¹ h⁻¹ with 98.41% selectivity.¹⁰⁹ This is very promising due to not only cost-effectiveness but also the outstanding performance. Despite these achievements,



Table 3 Photocatalytic dehydrogenation of C_2H_6 to C_2H_4

Photocatalyst	Cocatalyst	C_2H_4 production rate ($\mu\text{mol g}^{-1} \text{h}^{-1}$)	C_2H_4 selectivity	Reaction conditions	Ref.
TiO_2 (P25)	Pd	230.5	95.42%	300 W Xe lamp; 1 h; 25 mg catalyst; $CO_2: C_2H_6 = 1:1$; 0.2 MPa	108
TiO_2 (P25)	Cu	533.46	98.41%	UV lamp (CEL-HXF300); 1 h; 25 mg catalyst; Ar: $C_2H_6 = 9:1$; 0.2 MPa	109
$LaVO_4-O_v$	Pt	275	96.8%	300 W Xe lamp; 1 h; 100 mg catalysts; 150 mL pure C_2H_6	110
TiO_2 (P25)	Pd-Rh	428.8	68.7%	300 W Xe lamp; 1 h; 25 mg catalyst; $C_2H_6: CO_2 = 1:1$; 0.2 MPa	111
TiO_2 (P25)	Pd	614.9	94.6%	300 W Xe lamp; 1 h; 25 mg catalyst; $C_2H_6: CO_2 = 1:1$; 0.2 MPa	112
Black TiO_2	Pt	8.56	83.1%	300 W Xe lamp (equipped with 400 nm cut-off filter); 4 h; 50 mg catalyst; 0.1 mmol ethane	113
ZnO	PdZn	46.4 $\text{mmol g}^{-1} \text{h}^{-1}$	92.6%	100 W 365 nm LED lamp; 5 mg catalyst; reaction gas comprising C_2H_6 (5 vol% in Ar, 18 mL min^{-1}) and O_2 (1 vol% in Ar, 12 mL min^{-1}); Flow reactor	114

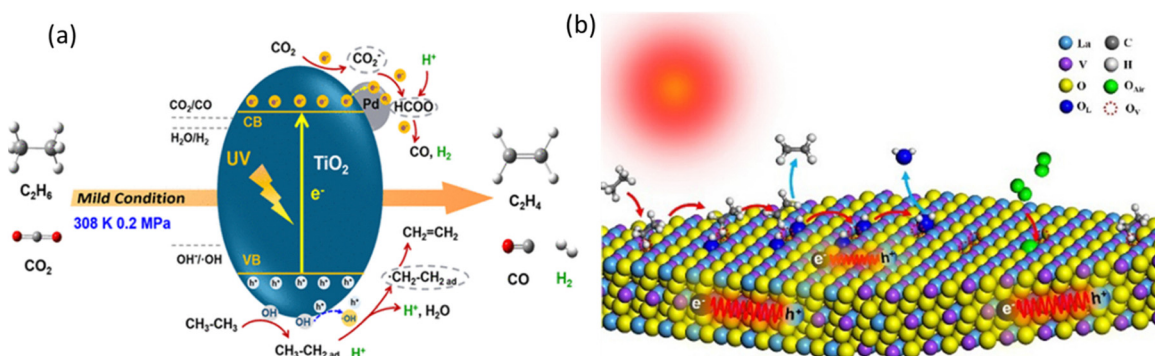


Fig. 17 (a) Schematic representation of (a) photocatalytic oxidative dehydrogenation of C_2H_6 with CO_2 over Pd/TiO_2 ; Reproduced with permission.¹⁰⁸ Copyright 2018, American Chemical Society and (b) photocatalytic C_2H_6 dehydrogenation over $LaVO_4-O_v$; reproduced with permission.¹¹⁰ Copyright 2024, Elsevier.

the field of photocatalytic C_2H_6 dehydrogenation to C_2H_4 is still in its infancy to construct the cascade $CH_4-C_2H_6-C_2H_4$ system *via* photocatalysis, particularly for non-noble metal-based cocatalyst design, which still requires exploration.

5 Conclusion and outlook

The increasing need for a scalable, one-step methane conversion process stems from environmental concerns and the growing demand for cleaner methods to utilize this abundant resource. The potential of EMC and PMC as viable techniques for the direct conversion of CH_4 into useful compounds such as oxygenates and hydrocarbons has been highlighted by this review. Current research in EMC encompasses both heterogeneous and homogeneous systems, offering alternative pathways for producing a range of products, including C_1-C_3 alcohols and carboxylic acids. Significant advancements in this area have been discussed, particularly strategies to mitigate side reactions associated with the oxygen evolution reaction (OER) through innovative catalyst and reaction system designs. These strategies include replacing active oxygen species from the OER with other sources, such as carbonate anions, oxygen gas, or *in situ* generated hydroxyl radicals. Notably, the OER-assisted EMC approach stands out as a highly promising method, leveraging the unique benefits of electrocatalysis. In this context, single-atom catalysts, such as Fe and FeNi composites,

represent cutting-edge advancements, though further research is needed to explore other electrocatalysts to enhance performance and efficiency.

PMC has also emerged as a promising approach, harnessing sunlight as an energy source. This review revisits the significant progress in CH_4 to C_2H_4 conversion *via* PMC, alongside strategies to enhance photocatalytic performance, including semiconductor catalyst design, cocatalyst loading, and reactor configuration. Addressing the remaining challenges requires continued research and innovation. Specifically, the development of low-cost, earth-abundant cocatalysts like Cu, Fe, Ni, and Co is essential to replace precious metals, making the process economically viable at an industrial scale. Additionally, the optimization of reactor configurations is crucial for improving mass transfer, light harvesting, and stability, warranting further attention in PMC research. Beyond the widely studied wide bandgap semiconductors like TiO_2 , WO_3 , and In_2O_3 , it is required to develop visible-light responsive semiconductors. A proposed photocatalytic $CH_4-C_2H_6-C_2H_4$ cascade system demonstrates promising performance for ethylene production, though further advancements in mass transfer, gas separation, and catalyst bed design are necessary. The field of photocatalytic methane coupling to ethylene holds significant potential for revolutionizing ethylene production through sustainable and eco-friendly methods. Continued interdisciplinary research and collaboration across chemistry, materials science, and engineering will be vital for translating these innovations into



practical, real-world applications, paving the way for a greener approach to methane conversion.

Data availability

No primary research results, software or code have been included and no new data were generated or analysed as part of this review.

Conflicts of interest

There are no conflicts to declare.

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