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# Advances in coupling catalytic selective oxidation reactions with *in situ* synthesis of hydrogen peroxide

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This review presents recent advances in coupling *in situ* hydrogen peroxide (H<sub>2</sub>O<sub>2</sub>) synthesis with selective oxidation reactions. As a green oxidant, H<sub>2</sub>O<sub>2</sub> plays an important role in the chemical industry. However, conventional production methods often yield highly concentrated H<sub>2</sub>O<sub>2</sub>, which is not suitable for direct use in reactions and raises significant safety concerns. The integration of *in situ* H<sub>2</sub>O<sub>2</sub> generation with selective oxidation allows for the immediate use of low-concentration H<sub>2</sub>O<sub>2</sub>, improving both safety and process efficiency. This review summarizes various strategies for *in situ* H<sub>2</sub>O<sub>2</sub> production, including enzymatic and catalytic approaches, and discusses their application in representative oxidation reactions such as olefin epoxidation, benzene hydroxylation, methane oxidation, adipic acid synthesis, Fenton processes, oxidative desulfurization, and the oxidation of sulfides to sulfones. Special attention is given to recent developments in catalyst composition and structural design, particularly in olefin oxidation. This review concludes with a summary of the advantages of *in situ* H<sub>2</sub>O<sub>2</sub> synthesis and offers perspectives on future research directions aimed at improving reaction efficiency, economic feasibility, and the development of sustainable green chemistry technologies.

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## 1 Introduction

Hydrogen peroxide (H<sub>2</sub>O<sub>2</sub>), an essential inorganic chemical raw material and a primary green chemical, has been listed as one of the top essential 100 chemicals globally. It has a high content of reactive oxygen species (mass fraction is 47%), and the degradation by-product is only water, eliminating the separation and purification process after the reaction. It has become a substitute for oxidants such as *t*-BuOOH, HNO<sub>3</sub>, N<sub>2</sub>O, and permanganate commonly used in industries. H<sub>2</sub>O<sub>2</sub> can be oxidized in a mild liquid environment, under acidic or alkaline conditions, compared with molecular oxygen. It is an efficient, strong multifunctional oxidant. Much research has been carried out to use H<sub>2</sub>O<sub>2</sub> as the oxidant in the

epoxidation of olefins, hydroxylation of aromatics, ammoxidation of ketones, and oxidation of alkanes. H<sub>2</sub>O<sub>2</sub> has excellent industrial application prospects due to its advantages of high selectivity, mild reaction conditions, and green environmental protection.<sup>1</sup> The industrial method of producing H<sub>2</sub>O<sub>2</sub> is mainly the anthraquinone method. The industrial technology of this method is mature, but there are some problems such as a large amount of equipment investment, complex process flow, and environmental pollution. Due to the limitation of process cost, H<sub>2</sub>O<sub>2</sub> production plants are often built in remote areas, which need to be stored and transported to the point of use for dilution in industrial applications. However, high-concentration H<sub>2</sub>O<sub>2</sub> is unstable, can easily decompose, has high storage and transportation cost, and has potential safety hazards. The direct synthesis of H<sub>2</sub>O<sub>2</sub> from H<sub>2</sub> and O<sub>2</sub> has become a research hotspot because of its advantages of high atomic economy and good environmental benefits. Furthermore, there has been considerable focus on both electrocatalytic<sup>2–4</sup> and photocatalytic<sup>5–7</sup> pathways for synthesizing H<sub>2</sub>O<sub>2</sub>. It is

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suitable for low-cost and small-scale production of  $\text{H}_2\text{O}_2$ . Prior to this, we also conducted related research on the direct synthesis of  $\text{H}_2\text{O}_2$  from hydrogen and oxygen.<sup>8,9</sup> The reaction mechanism is that  $\text{H}\cdot$ , which is formed by the dissociative adsorption of  $\text{H}_2$ , reacts with  $\text{OO}\cdot$  formed by non-dissociative adsorption of  $\text{O}_2$  to form  $\text{H}_2\text{O}_2$  through an  $\cdot\text{OOH}$  intermediate. There are many hydroperoxides and reactive oxygen species in the reaction

process, and the active species of  $\text{H}_2\text{O}_2$  act on the oxidation reaction. The following section introduces various strategies for *in situ*  $\text{H}_2\text{O}_2$  generation and, using representative oxidation reactions that are extensively studied both domestically and internationally as examples, reviews the recent progress and key challenges in this field, aiming to provide insights and references for future research.



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## 2 Approaches to *in situ* H<sub>2</sub>O<sub>2</sub> production

Driven by the need for greener and more sustainable chemical processes, the development of efficient methods for H<sub>2</sub>O<sub>2</sub> synthesis has become a research focus. Conventional anthraquinone-based processes suffer from complex procedures, high energy consumption, and undesired by-products, limiting their environmental viability. Akram *et al.* pioneered the direct synthesis of H<sub>2</sub>O<sub>2</sub> *via* the reaction of H<sub>2</sub> and O<sub>2</sub> over supported bimetallic gold–palladium catalysts. This innovative approach not only simplifies the production pathway but also demonstrates significant advantages in catalytic activity, selectivity, and stability, marking the inception of *in situ* synthesis methods for H<sub>2</sub>O<sub>2</sub> production.

### 2.1 Enzymatic catalysis method

Enzymatic catalysis represents a green and highly selective strategy for *in situ* H<sub>2</sub>O<sub>2</sub> generation. Typical systems employ redox enzymes such as glucose oxidase or peroxidases to catalyze the two-electron reduction of O<sub>2</sub> to H<sub>2</sub>O<sub>2</sub>, offering high substrate specificity and excellent biocompatibility.<sup>10</sup> Operating under mild conditions—ambient temperature and near-neutral pH—this method is particularly suitable for biologically or environmentally sensitive oxidation processes. The generation efficiency and selectivity of H<sub>2</sub>O<sub>2</sub> can be further enhanced by optimizing enzyme immobilization strategies, tuning electrode interfaces, and controlling electron transfer mechanisms, including both mediated and direct pathways.<sup>11</sup> Despite its potential, this approach is currently limited by enzyme instability, deactivation tendencies, and relatively low productivity.

### 2.2 Chemical catalysis method

Chemical catalytic synthesis of H<sub>2</sub>O<sub>2</sub> *via* the direct reaction of H<sub>2</sub> and O<sub>2</sub> over noble metal catalysts represents one of the most promising strategies for *in situ* H<sub>2</sub>O<sub>2</sub> generation. The mechanism involves dissociative adsorption of H<sub>2</sub> and non-dissociative adsorption of O<sub>2</sub> on the catalyst surface, leading to ·OOH intermediates that subsequently form H<sub>2</sub>O<sub>2</sub>. Suppressing the undesired hydrogenation of H<sub>2</sub>O<sub>2</sub> to water is key to improving selectivity.<sup>12</sup> Studies have shown that the catalyst's metal composition, oxidation state, support material, and reaction environment all significantly influence the activity and selectivity. Alloying,<sup>13</sup> the use of high-surface-area supports such as activated carbon, and fine control over gas–liquid–liquid interfaces in microreactor systems<sup>14</sup> have proven effective in enhancing both the efficiency and the process safety. Moreover, introducing metal or non-metallic additives can modulate the electronic properties of Pd, thereby suppressing the H<sub>2</sub>O<sub>2</sub> degradation pathways.<sup>15</sup> This approach offers high atom economy, a short reaction route, and strong compatibility with oxidation processes. However, challenges remain, including catalyst deactivation and the inherent safety risks associated with H<sub>2</sub>/O<sub>2</sub> mixtures.

### 2.3 Ultrasonic method

Ultrasound-assisted contact electro-catalysis has recently emerged as a novel and green strategy for *in situ* H<sub>2</sub>O<sub>2</sub> generation, offering advantages such as metal-free operation, environmental compatibility, and structural simplicity. Wang *et al.*<sup>16</sup> reported a catalyst-free system in which the combination of a poly(tetrafluoroethylene) stirring bar and ultrasound induces localized charge separation at the gas–liquid interface, facilitating the two-electron reduction of O<sub>2</sub> to H<sub>2</sub>O<sub>2</sub> under ultrasonic excitation, charge accumulation on the PTFE surface mimics the role of an electrode, enabling the formation of reactive oxygen species that subsequently convert into H<sub>2</sub>O<sub>2</sub>. This approach eliminates the need for noble metals or enzymatic components and operates under mild, low-cost conditions, making it attractive for sustainable oxidation applications. However, current limitations include relatively low H<sub>2</sub>O<sub>2</sub> production rates, inefficient interfacial charge separation, and limited system stability.

### 2.4 Plasma method

Plasma-assisted catalysis offers a non-thermal, high-energy alternative for *in situ* H<sub>2</sub>O<sub>2</sub> synthesis by activating molecular species through electron excitation. Zhou *et al.*<sup>17</sup> first demonstrated the direct synthesis of H<sub>2</sub>O<sub>2</sub> from H<sub>2</sub> and O<sub>2</sub> using non-equilibrium plasma discharge, in which reactive oxygen species, particularly ·OOH intermediates, are generated in the gas phase under plasma excitation. Subsequent work extended this concept to CH<sub>4</sub>/O<sub>2</sub> systems, where methane serves both as a stabilizer for plasma discharge and as a modulator of the H/O ratio, contributing to improved H<sub>2</sub>O<sub>2</sub> selectivity.<sup>18</sup> This plasma-based approach operates without traditional catalysts, offering advantages such as rapid start-up, broad operational flexibility, and compatibility with complex gas-phase mixtures. However, challenges remain, including relatively low H<sub>2</sub>O<sub>2</sub> yields, high energy consumption, and poor product stability due to harsh oxidative environments.

### 2.5 Photocatalytic method

Photocatalytic synthesis of H<sub>2</sub>O<sub>2</sub> involves the use of semiconductor materials to harness visible light for generating electron–hole pairs that drive the two-electron reduction of O<sub>2</sub>. Shiraishi *et al.*<sup>19</sup> reported that graphitic carbon nitride (g-C<sub>3</sub>N<sub>4</sub>) efficiently catalyzes this reaction under visible light, with high selectivity attributed to its electronic structure, which favors two-electron over four-electron O<sub>2</sub> reduction, thereby suppressing water formation. Building on this, Kofuji *et al.*<sup>20</sup> designed g-C<sub>3</sub>N<sub>4</sub>-based nanohybrids incorporating aromatic diimides and graphene, forming metal-free heterojunctions that promote charge separation and enable more stable photocatalytic H<sub>2</sub>O<sub>2</sub> production, achieving a solar-to-peroxide energy conversion efficiency of 0.2%. This strategy benefits from mild reaction conditions, solar energy utilization, and the absence of applied bias or precious metals, making it particularly



suitable for decentralized and green chemical systems. Nevertheless, limitations remain, including narrow visible light absorption ranges, high recombination rates of charge carriers, and challenges in stabilizing accumulated  $\text{H}_2\text{O}_2$ .

## 2.6 Electrochemical method

Electrocatalytic synthesis of  $\text{H}_2\text{O}_2$  via the two-electron oxygen reduction reaction represents a promising and sustainable route for *in situ*  $\text{H}_2\text{O}_2$  generation. This approach relies on applying a controlled potential to drive the selective reduction of  $\text{O}_2$  to  $\cdot\text{OOH}$  intermediates, avoiding the competing four-electron pathway to water, and thereby, enhancing the  $\text{H}_2\text{O}_2$  selectivity.<sup>21,22</sup> While early efforts focused on mechanistic understanding and precious metal catalysts,<sup>23</sup> recent advances emphasized the development of non-precious metal systems, particularly nitrogen-doped carbon-based materials and single-site M–N–C structures.<sup>24–26</sup> For instance, atomic-level tuning of coordination environments in Co–N–C catalysts had been shown to modulate the  $\text{O}_2$  adsorption geometry and balance the activity–selectivity trade-offs.<sup>25</sup> Electrocatalytic approaches offer several advantages including mild reaction conditions, high energy efficiency, and modular scalability for decentralized applications. However, challenges including catalyst degradation, limited Faradaic efficiency, and instability of  $\text{H}_2\text{O}_2$  in aqueous media remain.

## 2.7 Mechanochemical method

Mechanocatalysis has emerged as a novel strategy for *in situ*  $\text{H}_2\text{O}_2$  generation, driven by mechanical force-induced charge separation without the need for external light, heat, or electrical input. Jin *et al.*<sup>27</sup> demonstrated that ferroelectric  $\text{K}\text{Sr}_2\text{Nb}_3\text{Ta}_2\text{O}_{15}$  nanorods can produce surface potential under mechanical agitation, enabling the two-electron reduction of  $\text{O}_2$  to  $\text{H}_2\text{O}_2$  under ambient, dark conditions without any co-catalysts or redox mediators. This method features an energy-free, additive-free system with minimal infrastructure requirements, offering potential for decentralized green oxidation applications. However, current limitations include low  $\text{H}_2\text{O}_2$  productivity, challenges in controlling the reaction kinetics, and the need to improve the mechanical durability of the catalytic materials. Further mechanistic investigations are required to establish the structure–activity relationships and scalability of this approach.

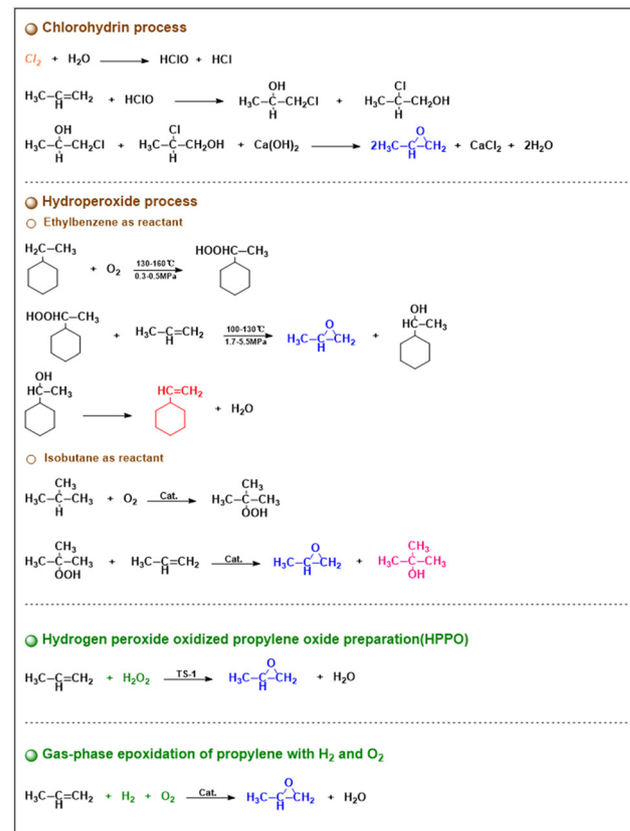
# 3 Propylene epoxidation to propylene oxide

## 3.1 Background

Propylene oxide (PO), a crucial intermediate in chemicals, is widely used to produce polyether polyols, propene glycols, and polyurethane, which are subsequently applied to manufacture polyurethane foams and polyesters. The current annual PO production is more than 10 million tons

worldwide, and the demand is increasing remarkably (Scheme 1).<sup>28,29</sup>

The current PO production processes include the chlorohydrin process and the hydroperoxide process (HPPO). Besides, Dong and coworkers<sup>30</sup> explored the epoxidation of olefins by an electro-organic system. Wang *et al.*<sup>31</sup> and Ko *et al.*<sup>32</sup> studied the epoxidation of propylene in a photocatalytic system and a photo-electro-heterogeneous catalytic system. Direct hydrogen peroxide synthesis offers distinct advantages over electrochemical and photocatalytic methods. It typically follows a simple, efficient reaction pathway under ambient temperature and pressure, eliminating the need for high voltages in electrochemical processes or light sources in photocatalysis. This improves energy efficiency and simplifies operations. The catalysts used in direct synthesis show high selectivity, minimizing side reactions, and demonstrate excellent stability, maintaining activity over multiple cycles. Additionally, the process does not require external power or specialized lighting, reducing system complexity and operational costs. However, the chlorohydrin process in the industry produces harmful by-products such as waste salt. As people pay more attention to the environment, the chlorohydrin process which produces harmful by-products is gradually eliminated. Although the hydroperoxide process avoids environmental



**Scheme 1** Various production routes of propylene oxide (PO). Reproduced from ref. 33 with permission from Elsevier, copyright 2021.



pollution, it is not cost-effective due to the lengthy process, high raw material requirements, and many by-products. It is urgent to find alternative production solutions.

H<sub>2</sub>O<sub>2</sub>, the oxidant used in the direct epoxidation of propylene (C<sub>3</sub>H<sub>6</sub>) to produce PO (HPPO), is more environmentally friendly, and the only by-product is water. Recently, researchers have obtained a deep understanding of the mechanism of this reaction and have scaled it up to industrial production. However, due to the high cost and risk of transporting H<sub>2</sub>O<sub>2</sub>, an H<sub>2</sub>O<sub>2</sub> manufacturing plant is usually built close to the PO plant to obtain the highest economic benefit. At present, considerable research efforts are dedicated to synthesizing H<sub>2</sub>O<sub>2</sub> by H<sub>2</sub> and O<sub>2</sub> first and then consuming it *in situ* to oxidize propylene to PO. Compared with the HPPO process, this gas–solid reaction shows obvious advantages of easy operation, simple product separation, and high economic benefits.<sup>33</sup>

Due to the unique synergy between the Au site and the Ti site, the current research mainly focuses on applying titanium-containing materials loaded with gold nanoparticles as catalysts for the epoxidation of propylene with hydrogen and oxygen. In addition, some researchers try to use Ni (ref. 34 and 35) as an active metal or ZrO<sub>2</sub> (ref. 36) as a support to provide some new strategies. Here, we mainly review the influence mechanism of the structure of gold supported over Ti-based materials as the catalyst on reaction performance and the strategies to improve the catalytic performance in *in situ* propylene epoxidation systems in recent years, such as the influence of metal particle sizes, deposition methods, support structure, and support surface groups on reaction performance.

### 3.2 Effect of metal

**3.2.1 Gold particle sizes.** Gold nanoparticles of different sizes have been used in various reactions. Single-atom, gold bilayer, sub nanoclusters, clusters (1–2 nm), and NPs (2–5 nm) were loaded onto different supports and found to have unique characteristics.<sup>37</sup> In the epoxidation of propylene with hydrogen and oxygen, the size effect of Au particles is one of the research focuses. Accurately revealing the effect of gold particle size on the reaction will be helpful to the design of gold catalyst.<sup>38</sup>

Some researchers believe that the size of gold particles should be less than 2.0 nm in the reaction system of *in situ* epoxidation of propylene with hydrogen and oxygen. Qin Zhong and coworkers<sup>39</sup> considered that Au nanoclusters (<1.0 nm) are the primary active sites of gold for propylene epoxidation. Three different supports (titanium silicalite-1 (TS-1), silicalite-1 (S-1)/TS-1, and uncalcined titanium silicalite-1 (unTS-1)) were prepared using the hydrothermal method, and gold was loaded using the DP method. It has been found that the gold nanoclusters (<1.0 nm) on the external surfaces of TS-1 and inside the nanoporous channels of TS-1 are the dominant gold active sites, which is due to its unique catalytic properties. First, gold exhibits high surface

energy and abundant active sites at the nanoscale, which effectively activate oxygen molecules to form superoxide species (O<sub>2</sub><sup>-</sup>), thereby promoting the selective oxidation pathway of the reaction. Second, gold possesses a weak adsorption ability for C–H bonds, which helps to prevent the side reactions (*e.g.*, deep oxidation) and enhances the selectivity for propylene oxide. Additionally, the surface electronic properties of gold can synergize with supports (such as TiO<sub>2</sub> or SiO<sub>2</sub>), further improving the catalytic activity and stability. These characteristics enable gold to achieve both high activity and high selectivity in the epoxidation of propylene, making it the optimal catalyst. In addition, they hope to prepare smaller gold nanoparticles to obtain better catalytic performance. Cheng and coworkers<sup>40</sup> prepared a series of Au/Si/Ti-MCM-36 and Au/Ti-YNU-1 with varying gold loading. They found that the Au nanocluster size of the most efficient catalyst in this series is 1.04 nm. The highly isolated TdTi in Ti-YNU-1 favors the formation of smaller gold nanoclusters.

However, other researchers believe that the optimized size of gold particles should be 2.0–5.0 nm. Feng and coworker<sup>41</sup> found that as the size of the Au particles increased from 2.6 nm to 5.1 nm, the PO selectivity decreased monotonically from 90% to 61%. At the same time, the by-products (such as carbon dioxide, propionaldehyde, ethanol, and acetone) are increasing. Recent advances in nanostructured catalysts demonstrate that confining Au clusters within two-dimensional MXene layers (Ti<sub>3</sub>C<sub>2</sub>T<sub>x</sub>) significantly enhances stability and selectivity. For instance, Li and workers designed Au/MXene-TS-1 composites where the MXene's interlayer spacing (≈1.2 nm) restricts Au growth to 2–3 nm clusters. This confinement not only prevents sintering but also facilitates electron transfer from Ti sites to Au, achieving 97% PO selectivity at 200 °C with a record turnover frequency (TOF) of 1200 h<sup>-1</sup>.<sup>42</sup> *In situ* TEM studies further revealed that MXene's oxygen-terminated surface stabilizes Au<sup>δ-</sup> species, which are critical for H<sub>2</sub>O<sub>2</sub> activation. To study the size effect of supported Au more effectively, Huang and coworkers<sup>43</sup> loaded Au NPs/clusters on nonporous and highly crystalline TiO<sub>2</sub>- surface.<sup>44</sup> *Via* a comprehensive *in situ* DRIFTS study, they found that Au particles with sizes of 1.4 to 4.6 nm are mainly composed of the Au<sup>δ-</sup> species. An increase in the Au particle size to 5.0 nm increases the active Au<sup>δ-</sup>-Ti<sup>4+</sup> ensembles, further increasing the Au particle size to 7.0 nm results in a substantial increase in the Au<sup>δ+</sup> species. Li and coworkers<sup>45</sup> synthesized TS-1 *via* a seed-induced method, and bio-inspired Au nanoparticles were loaded through an ionic-liquid-enhanced-immobilization method onto the TS-1 support. They demonstrated that the dosage of the seed introduced during the synthesis of TS-1 significantly affects the acidity of the resulting TS-1, which plays an essential role in regulating the size of the Au NPs (3.0–4.1 nm), and the size of Au nanoparticles within the range of 3.2–3.9 nm was found to be optimal for propylene epoxidation.

The most suitable gold particle size is still controversial, which may be related to supports and deposition methods.





revealed negligible deactivation, which is attributed to the ability of CeO<sub>2</sub> to scavenge reactive oxygen species (ROS), thereby mitigating catalyst degradation. Notably, each promoter exhibits distinct functionalities and plays a unique role in enhancing the overall catalytic efficiency and durability of the system.

### 3.3 Effect of support

Many titanium-containing materials such as TiO<sub>2</sub>, TiO<sub>2</sub>-SiO<sub>2</sub>, Ti-SiO<sub>2</sub>, TS-1, and TS-2 have been used in propylene epoxidation with hydrogen and oxygen. In recent years, the research on TS-1 has increased significantly due to its isolated Ti sites, stable structure, rich pores, and other excellent properties. Research on TS-1 mainly focuses on adjusting its structure and surface groups. The PO formation rates of different catalysts are shown in Scheme 2.

**3.3.1 Pore structure of supports.** In recent years, many researchers have paid attention to the effect of support pore structure on catalytic performance, and many titanium silicon molecular sieves such as mesoporous titanium silicate-1 (MTS-1),<sup>56,57</sup> small-sized mesoporous titanium silicate-1 (STS-1)<sup>58</sup> and hierarchical titanium silicate-1 (HTS-1)<sup>59,60</sup> have been used to improve the performance of catalysts. The stability and mass transferability of these carriers have been widely recognized. The better mass transferability by shortened reactant/product diffusion length results in lower coke formation and the absence of refractory aromatic coke, inhibiting the side reactions and deactivation caused by the blocking of micropores and active sites. For example, Zhuang *et al.*<sup>61,62</sup> introduced Fe single-atoms into hierarchical TS-1 (Fe-SA/TS-1) by a vapor-phase exchange method. The Fe-N<sub>4</sub> sites adjacent to Ti-O<sup>-</sup> centers act as co-catalysts, lowering the activation energy for H<sub>2</sub>O<sub>2</sub> formation from 85 kJ mol<sup>-1</sup> to 62 kJ mol<sup>-1</sup>. Coupled with the hierarchical pores (micropores: 0.55 nm; mesopores: 5–10 nm), this design achieved 90% H<sub>2</sub>O<sub>2</sub> utilization efficiency—40% higher than pure TS-1—by ensuring rapid diffusion of H<sub>2</sub>O<sub>2</sub> to Au active sites while minimizing decomposition.

**3.3.2 Supports with core-shell structures.** The influence of the core-shell structure of the support on the catalytic performance has also attracted much attention in recent years. Many novel catalysts have been designed and applied in the catalyst system of *in situ* epoxidation of propylene with hydrogen and oxygen, such as Au/S-1/TS-1,<sup>39,63</sup> Au/TS-1/S-1,<sup>64</sup> Au/TS-1@S-1,<sup>65</sup> Au-Ti@MFI,<sup>66</sup> and Au/S-1/TS-1@dendritic-SiO<sub>2</sub>.<sup>67</sup> These catalysts with core-shell structures have dispersed active sites, excellent mass transferability, and high stability, and some unique sites often appear at the core-shell junction.

**3.3.3 Effect of support surface groups.** The groups on the surface of the support will also affect the performance of the catalyst. Angelo and coworkers<sup>68</sup> found that hexamethyldisilazane (HMDS) and transition metal

dichalcogenides (TMDs) for gas-phase silylation can significantly improve the catalytic performance. Hexamethylenediamine (HMD) improved the yield and selectivity of PO, while TMD treatment completely inhibited the formation of propane. These findings show that the silylation of the catalyst can provide hydrophobicity for the catalyst and promote the desorption of the product but also effectively block the sites of side reactions.

The schematic diagram of catalyst design in propylene epoxidation and experimental parameters and results of different researches are shown in Table 1 and Scheme 3. In the direct gas-phase epoxidation of propylene, Au nanoparticles and titanium-containing supports form a series of bifunctional catalysts. TS-1 has excellent catalytic performance among many titanium-containing supports because of its isolated tetra-coordinated Ti and strong hydrophobicity. In recent years, researchers have carried out comprehensive research and exploration on such catalysts to improve further the catalytic performance and stability of catalysts from the perspectives of gold particle-size regulation and loading, the addition of promoters, pore structure support, core-shell structure, and surface modification. At the same time, they also provide strategies and methods for the follow-up research of this reaction and similar reactions.

### 3.4 Selective oxidation of ethylene to ethylene glycol *via in situ* H<sub>2</sub>O<sub>2</sub>

Among diverse olefin oxidation reactions, the selective oxidation of ethylene to ethylene glycol (EG) represents a transformation of considerable industrial significance. As the simplest olefin and a cornerstone petrochemical feedstock, ethylene derivatives are foundational to numerous industrial sectors, making EG production a key indicator of chemical manufacturing capacity.<sup>87,88</sup> Ethylene glycol, the simplest aliphatic diol, is widely used in the synthesis of polyester fibers, plastics, and related materials, and serves as a critical benchmark for assessing industrial chemical development.<sup>89,90</sup>

The choice of oxidant fundamentally governs the efficiency of ethylene-to-EG processes. While molecular oxygen (O<sub>2</sub>) is commonly used, its high reactivity often results in the overoxidation to CO<sub>2</sub> and water, complicating downstream separation.<sup>91</sup> Air, though more economical, suffers from lower oxygen content (~21%) and correspondingly reduced selectivity. Process data typically show an EG selectivity of 65–75% when using air, compared to 75–90% with pure O<sub>2</sub>, with ethylene conversion rates generally around 8–12%.<sup>92</sup> In contrast, peroxide-based oxidants, particularly hydrogen peroxide (H<sub>2</sub>O<sub>2</sub>), offer significant advantages by enabling up to 100% selectivity to EG with minimal side-product formation.<sup>93</sup> Furthermore, H<sub>2</sub>O<sub>2</sub> decomposes into only water and oxygen, aligning with the principles of green chemistry.

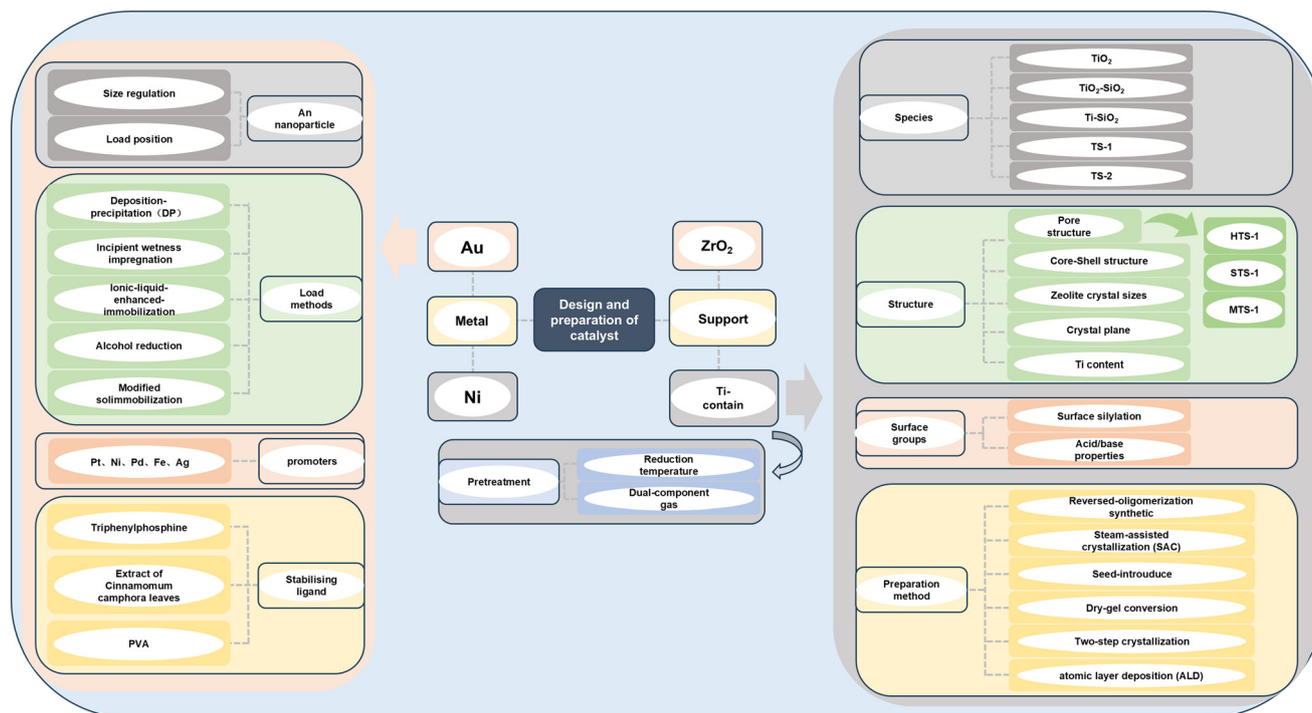
Guan *et al.*<sup>94</sup> established an integrated electrocatalytic tandem system for ethylene-to-EG conversion, employing



Table 1 Summary of various catalytic systems for coupling the epoxidation of propylene with *in situ* generation of H<sub>2</sub>O<sub>2</sub>

Ref.	Catalyst		Experimental conditions					Activity results				
	Catalyst name	Au loading (wt%)	Au particle size (nm)	Catalyst mass (g)	T (K)	GHSV (ml g <sub>cat</sub> <sup>-1</sup> h <sup>-1</sup> )	C <sub>3</sub> H <sub>6</sub> :H <sub>2</sub> :O <sub>2</sub> :inert	Propylene conversion (mol%)	PO selectivity (mol%)	H <sub>2</sub> efficiency (mol%)	Reaction rate (mmol <sub>PO</sub> h <sup>-1</sup> g <sub>Au</sub> <sup>-1</sup> )	(g <sub>PO</sub> h <sup>-1</sup> kg <sub>cat</sub> <sup>-1</sup> )
43	Au/TiO <sub>2</sub>	5	5 ± 1.6	0.25	323.15	8000	1:1:1:7	1.8	96.8	33	12.52	36.3
69	Au/Ti-SiO <sub>2</sub>	0.18	2.3 ± 1.0	0.15	473.15	10000	1:1:1:7	2.2	89.3	4.8	396.55	41.4
68	Au/Ti-SiO <sub>2</sub>	0.15	—	0.15	493.15	10000	1:1:1:7	4.7	83	11	1160.92	101
70	Au/Ti-SiO <sub>2</sub>	0.03	2.4 ± 0.6	0.15	473.15	14000	1:1:1:7	0.8	84.9	12.8	1390.80	24.2
71	Au/TS-1	0.24	—	0.15	473.15	14000	1:1:1:7	12	87	25	2693.97	375
72	Au/TS-1	0.314	5.1 ± 4.7	0.15	473.15	14000	1:1:3:5	—	98	26	1921.81	350
39	Au/TS-1	0.1	—	0.15	473.15	14000	1:1:1:7	10.5	87	10	5689.66	330
73	Au/TS-1	2	3.7	0.1	473.15	9000	1:1:1:7	5.2	84.83	25	88.74	102.94
74	Au/TS-1	0.15	2.1 ± 0.3	0.15	473.15	8000	1:1:1:17	8.4	81.3	14.2	765.52	66.6
45	Au/TS-1	—	3.2 ± 0.5	0.15	578.15	7000	1:1:1:7	19	70.5	72	—	221
29	Au/TS-1	1	3.0 ± 0.8	0.5	473.15	8000	1:1:1:7	1.3	88	23	38.79	22.5
49	Au/TS-1	0.1	2.4 ± 0.5	0.15	473.15	14000	1:1:1:7	—	83	—	3534.48	205
54	Au-Pd/TS-1	0.76	3.9 ± 0.6	0.15	473.15	14000	1:1:1:7	—	42	—	90.74	40
51	Au-Fe/TS-1	0.69	3.1 ± 0.2	0.15	473.15	14000	1:1:1:7	7.3	89.9	25.7	596.70	238.8
46	AuNi/TS-1	0.212	3.5 ± 0.6	0.15	443.15	14000	1:1:1:7	10.3	73	31	1252.44	154
75	Au/TS-1-B	0.09	2.8 ± 0.5	0.15	473.15	14000	1:1:1:7	—	91	40	2758.62	144
76	Au/TS-1-B	0.5	4.6 ± 0.5	0.15	578.15	7000	1:1:1:7	14.5	73.7	28	644.83	187
77	Au/TS-1-B	—	3.1 ± 0.6	0.15	473.15	14000	1:1:1:7	—	88	37	—	150
41	Au/TS-1-B	0.13	2.6 ± 0.5	0.15	473.15	14000	1:1:1:7	9	70	15	2917.77	220
52	AuAg/TS-1-B	0.08	2.8 ± 0.5	0.15	493.15	14000	1:1:1:7	—	73	44	5021.55	233
78	AuPt/TS-1-B	0.1	3.2 ± 0.7	0.15	473.15	14000	1:1:1:7	11	88	41	6137.93	356
50	Au/TS-1-SA	0.15	2.8 ± 0.5	0.15	473.15	14000	1:1:1:7	—	90	15	1758.62	153
79	Au/TS-1-S5	0.5	3.8 ± 0.6	0.15	573.15	7000	1:1:1:7	18.3	72.6	32.2	757.59	219.7
80	Au/TS-1-NaC-CsC	0.21	2.7 ± 0.9	0.15	473.15	8000	1:1:1:17	9.2	91.2	18.1	665.02	81
81	Au/TS-1-NaC	0.25	1.9 ± 0.5	0.1	483.15	14000	1:1:1:55.5	—	95	—	151.72	22
82	Au/TS-2-B	0.09	2.9 ± 0.4	0.15	473.15	—	1:1:1:7	—	91	35	2260.54	118
59	Au/HTS-1	0.1	3.2 ± 0.2	0.15	473.15	—	1:1:1:7	—	85	—	2155.17	125
60	Au/HTS-1	0.1	3.0 ± 0.3	0.15	473.15	14000	1:1:1:7	—	90	—	2586.21	150
57	Au/MTS-1	0.13	3.4 ± 0.5	0.15	473.15	14000	1:1:1:7	—	95.2	—	1883.29	142
56	Au/MTS-1	0.324	—	0.15	473.15	14000	1:1:1:7	—	95	—	1543.21	290
83	Au/PTS-1	0.17	—	0.15	473.15	14000	1:1:1:7	6.8	91	—	1926.98	190
58	Au/STS-1	0.1	3.1 ± 0.4	0.15	473.15	—	1:1:1:7	—	91.2	—	2431.03	141
84	Au/UTS-1	0.031	—	0.15	473.15	14000	1:1:1:7	1.08	97	24	3170.19	57
85	Au/R-TS-1	—	1.98	0.15	473.15	14000	1:1:1:7	—	86	20	—	320
86	TiO <sub>2</sub> /Au/SiO <sub>2</sub>	0.7	—	0.15	373.15	14000	1:1:1:7	—	90	—	3.69	1.5
40	Au/A-Ti-YNU-1	0.13	1.5 ± 0.5	0.25	498.15	7000	1:1:1:7	3.3	59.3	36.2	397.88	30
48	Au/mesoTi-O-Si-50	0.5	2.4 ± 0.3	0.15	573.15	7000	1:1:1:7	12.8	76.2	18.2	554.83	160.9
64	Au/TS-1/S-1	0.1	2.9 ± 0.3	0.15	473.15	14000	1:1:1:7	—	87.2	—	2413.79	140
63	Au/S-1/TS-1	0.326	—	0.15	473.15	14000	1:1:1:7	6.7	96	15	2168.39	410
65	Au/TS-1@S-1	0.326	4	0.15	473.15	—	1:1:1:7	—	—	—	105.00	350
67	Au/S-1/TS-1@dentritic-SiO <sub>2</sub>	0.1	2.9 ± 0.3	0.15	473.15	14000	1:1:1:7	4.18	93.9	26.1	2472.41	143.4
66	Au-Ti@MFI	0.5	—	0.15	573.15	7000	1:1:1:7	14.8	83.9	—	706.90	205
34	Ni/Ti-SiO <sub>2</sub>	1	3.2 ± 0.4	—	473.15	10000	1:1:1:7	7.5	90.7	16.3	—	112
36	Au/ZrO <sub>2</sub>	0.017	3.6 ± 0.6	0.15	423.15	8000	1:1:1:7	0.14	42.3	—	124.40	122.65





**Scheme 3** Design framework of the propylene epoxidation catalyst.

electrocatalytic oxygen reduction on mesoporous carbon (MC-3) to generate  $\text{H}_2\text{O}_2$  *in situ*, followed by immediate ethylene oxidation over TS-1 zeolite. Their approach achieved 99% EG selectivity and 86%  $\text{H}_2\text{O}_2$  conversion under ambient temperature and pressure, effectively circumventing the high energy demands of conventional processes and eliminating the risks associated with  $\text{H}_2\text{O}_2$  storage and transport. The use of noble-metal-free catalysts further highlights the sustainability and scalability of this method.

In addition, Yang *et al.*<sup>95</sup> engineered a vanadyl phthalocyanine-functionalized catalyst (VOPc/CNT-OH) for highly selective  $\text{H}_2\text{O}_2$  production in acidic media. Industrial feasibility was demonstrated through continuous operation exceeding 100 hours in flow cell and solid-state electrolyte (SSE) reactors. The electrogenerated  $\text{H}_2\text{O}_2$  was successfully utilized for ethylene oxidation to EG, validating the practicality of coupled electrocatalytic–thermocatalytic systems.

Consequently, *in situ* reaction coupling strategies offer compelling advantages, including process simplification, the elimination of  $\text{H}_2\text{O}_2$  storage and transportation hazards, and improved economic viability. These developments demonstrate significant potential for industrial implementation in sustainable EG production.

## 4 Hydroxylation of benzene to phenol

Phenol is an essential intermediate product widely used in synthesizing resins, drugs, dyes, and other fields. The global phenol market has been valued at over \$20 billion in recent

years and is expected to maintain a considerable growth rate before 2025.<sup>96</sup> Over 90% of the phenol synthesized today is made by the three-step cumene process, which includes a high-energy condition. Initially, the cumene produced through the alkylation of benzene with propylene is oxidized to cumene hydroperoxide. Then, the phenol and by-product acetone are produced through the hydrolysis of cumene hydroperoxide. The cumene process has a low phenol yield and suffers from low atomic utilization, environmental unfriendliness, high energy consumption, multiple steps, and large amounts of by-products. This method faces high by-product separation costs when separating the by-product acetone.

Furthermore, the risk of explosion potentially exists in the cumene process. To address these challenges, an alternative method called direct hydroxylation of benzene to phenol (DHBP) has been proposed. This direct hydroxylation of benzene remains one of the most challenging catalytic reactions since the benzene ring must be broken in this process.<sup>97</sup>

### 4.1 Direct hydroxylation of benzene to phenol

$\text{O}_2$ ,  $\text{N}_2\text{O}$ , and  $\text{H}_2\text{O}_2$  are the common oxidants utilized in the DHBP process due to their exceptional oxidation capabilities. As early as 1983, Iwamoto *et al.*<sup>98</sup> reported the application of  $\text{N}_2\text{O}$  on  $\text{V}_2\text{O}_5\text{-SiO}_2$  catalysts, achieving 11% conversion of benzene and 45% yield of phenol. Although the unique oxidation effects are obtained by adopting  $\text{N}_2\text{O}$  (ref. 99 and 100) and  $\text{H}_2\text{O}_2$ ,<sup>101–103</sup> their practical applications are limited



due to their operating condition and material costs. However, DHBP is available by using O<sub>2</sub> as an oxidant instead. The direct synthesis of phenol from benzene and oxygen requires the activation of the C–H bond in the aromatic ring and the subsequent insertion of oxygen. The oxidation of benzene *via* an electrophilic reaction requires the oxygen species with a negative charge provided in most heterogeneous catalysts containing transition metals such as O<sup>-</sup>, O<sup>2-</sup>, and O<sup>2-</sup>.<sup>104</sup>

Because of the considerable economic and environmental advantages of this method, direct oxidation of benzene to phenol by using a mixture of H<sub>2</sub> and O<sub>2</sub> has been widely studied. However, many reports show high phenol selectivity using gaseous H<sub>2</sub> and O<sub>2</sub> as reactants but still low phenol yields, or very low phenol selectivity but relatively high yields. Besides, due to a mixture of H<sub>2</sub> and O<sub>2</sub>, the risk of explosion should be considered seriously.

#### 4.2 Pd membrane reactor

In this system, H<sub>2</sub> and O<sub>2</sub> are supplied separately on each side of the membrane to avoid contact in the gas phase. In addition, only H<sub>2</sub> can pass through the dense Pd membrane, while O<sub>2</sub> cannot. *In situ* H<sub>2</sub>O<sub>2</sub> and other reactive oxygen species (such as HOO· and HO·), which are able to promote the conversion of benzene to phenol, are generated while oxygen is reacting with the dissociated hydrogen that permeates out of the one side of the Pd membrane.<sup>105–108</sup> Thus, this kind of Pd membrane reactor is considered an economic, clean, and valuable technology.

The palladium membranes can be prepared by chemical vapor deposition (CVD) or electroless plating (ELP). CVD is suitable for industrial applications requiring high-quality, complex structures, and large-area films, while ELP is more suited for small-scale production, where the membrane quality is less critical. Noteworthy, the Pd membrane is not only a catalyst but also a hydrogen separator. In contrast to conventional Pd-loaded catalysts in porous supports, the surface area of the Pd membrane is relatively small. The application of Pd membranes in direct catalytic hydroxylation of benzene to phenol is rapidly gaining plenty of attention after Niwa reported this kind of membrane reactor. However, the phenol yields of many studies are not satisfactory.<sup>109–118</sup> Zhang *et al.*<sup>119</sup> employed operando X-ray absorption spectroscopy (XAS) and surface-enhanced Raman spectroscopy (SERS) to track the dynamic evolution of Pd species during the reaction. They identified a reversible Pd<sup>0</sup>

↔ PdO<sub>x</sub> transformation cycle, where Pd<sup>0</sup> domains (2–3 nm) facilitate H<sub>2</sub> dissociation, while PdO<sub>x</sub> interfaces (PdO/PdO<sub>2</sub>) stabilize HOO· intermediates. By precisely controlling the H<sub>2</sub>/O<sub>2</sub> ratio at 2/1, the Pd<sup>0</sup>/PdO<sub>x</sub> ratio was optimized to 3/1, yielding 22.3% phenol with 98% selectivity—a 50% improvement over previous membrane reactors. Vulpescu *et al.*<sup>118</sup> pointed out that commercial application of Pd membranes is possible through computer simulations if the productivity of phenol and the low efficiency of raw materials are solved. The formation of many by-products was demonstrated to occur *via* two significant side reactions that dominate under the applied experimental conditions. These two reactions are the complete oxidation of hydrogen to water and the complete oxidation of benzene to carbon dioxide. Besides, Niwa *et al.*<sup>105</sup> found membrane deterioration by releasing the Pd layer from the alumina support tube, and Vulpescu *et al.*<sup>118</sup> and Shu *et al.*<sup>114</sup> reported that hot spots can lead to membrane deterioration under oxygen-enriched conditions. Non-selective oxidation and complete combustion occur in a higher oxygen content. However, a higher hydrogen content favors hydrogenation reactions.<sup>117,118,120</sup> Moreover, Sato *et al.*<sup>104</sup> indicated that two phases of palladium hydride with different cell sizes coexist, that is, α-phase and β-phase at below 566.15 K, leading to embrittlement of the Pd membrane. Currently, outstanding results were reported by Wang *et al.*<sup>97,106,121–123</sup> through a systematic study of the membrane, parameters, and feed mode (Table 2).

Even though the Pd membrane reactor has shown encouraging results, space for further commercialization improvement exists due to weaknesses such as membrane deterioration and hot spots, poor membrane stability, and low feedstock utilization.

## 5 Selective oxidation of methane into oxygenates

The resources of methane are diverse. The main existing form of methane is natural gas, and it makes up nearly 90% of natural gas.<sup>124</sup> The reserves of natural gas are abundant on the earth. It is plentiful in the crust around the world, constituting approximately 21% of the total principal energy sources on earth.<sup>125</sup> The development of simple processes for upgrading natural gas to value-added chemicals is therefore highly desirable.<sup>126</sup> The functionalization of methane to valuable hydrocarbons and oxygen-containing products is

**Table 2** Comparison of different results about DBHP over Pd-based membrane reactors

Ref.	Membrane system	Method	T (K)	Benzene conversion (%)	Phenol selectivity (%)	Phenol yield (%)	Long-term (h)
106	Pd-TS-1 membrane	ELP	473.15	7.3	95	6.9	100
106	Pd capillary membrane	ELP	473.15	19	86	16.3	100
123	Pd capillary membrane	ELP	523.15	22.4	76	17.1	140
122	Pd-TS-1p membranes	ELP	473.15	7.2	99	7.1	na.
97	Pd-TS capillary membrane	ELP	473.15	23.4	95	22.3	110
121	Pd-TSH membrane	ELP	473.15	21.9	98	21.4	120



meaningful. Mainly, selective oxidation of methane to methanol is one such process that has received significant attention.<sup>127</sup>

Methane ( $\text{CH}_4$ ) is the simplest saturated hydrocarbon with a standardized tetrahedral structure. The strong C–H bond ( $104 \text{ kcal mol}^{-1}$ ,  $1 \text{ cal} = 4.18 \text{ J}$ ) with negligible electron affinity causes difficulty in C–H activation under mild conditions. The methanol product is also more active than methane and is more prone to overoxidation. Consequently, selective methane oxidation to methanol under mild conditions has been considered a dream reaction (Scheme 4).<sup>128</sup>

Conventionally, an indirect route for converting natural gas to methanol is used. The process has two steps: (1) intermediate production of synthesis gas by steam reforming and (2) catalytic conversion of synthesis gas to methanol. However, syngas production is an energy-intensive process, operated at 65% thermodynamic efficiency between 1073.15 and 1273.15 K, and more than 25% of the feed (natural gas) has to be burned to provide the heat for reaction. There are drawbacks such as complex process flows, low raw material utilization, high carbon emissions, and short catalyst lifespan. Hence, from the viewpoint of sustainable development, the direct conversion of methane to methanol is a desired alternative to the current technology.<sup>124</sup>

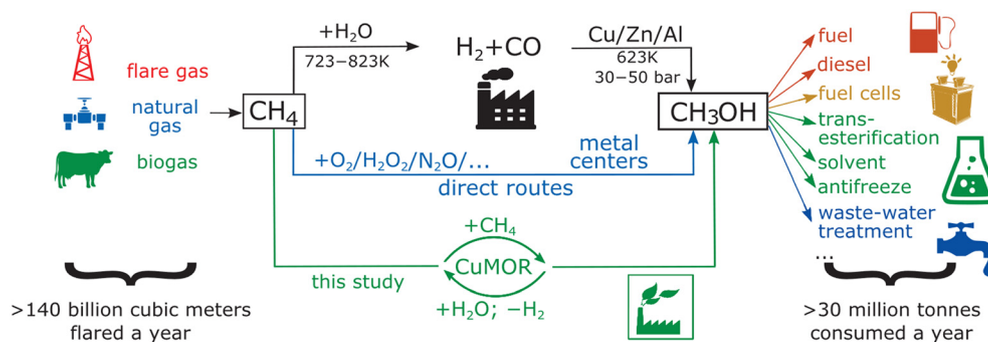
In order to achieve the direct oxidation of methane to methanol, several relatively expensive and toxic oxidants have been employed with catalysts, such as fuming sulfuric acid with Hg or Pt catalysts and selenic acid with Au and Pt catalysts. This method could produce methanol and overoxides and environmentally harmful by-products. The metal-exchanged zeolite catalysts could catalyze the methane oxidation using  $\text{N}_2\text{O}$ ,  $\text{O}_2$ , or even  $\text{H}_2\text{O}$ , which efficiently inhibited the methanol overoxidation, but still required a reaction temperature of  $200 \text{ }^\circ\text{C}$  or higher.<sup>128</sup>  $\text{H}_2\text{O}_2$ , an environmentally friendly oxidant, has significant advantages in methane oxidation under mild conditions without any toxic salts or strong acids. Using  $\text{H}_2\text{O}_2$  as an oxidant in the direct methane-to-methanol conversion holds potential for improved efficiency and reduced environmental impact, though challenges remain in catalyst stability, cost, and scaling up for industrial production. However,  $\text{H}_2\text{O}_2$  is

expensive relative to gaseous oxygen. In addition,  $\text{H}_2\text{O}_2$  in the presence of  $\text{O}_2$  has been used in the selective oxidation of methane under mild conditions to lower the production costs.<sup>129</sup>

For further control of the cost of this reaction, recent exploration was performed in combining the selective oxidation of methane with the *in situ* generation of  $\text{H}_2\text{O}_2$  using  $\text{H}_2$  and  $\text{O}_2$  (Table 3).

Hutchings and co-workers<sup>130–134</sup> have demonstrated that titania-supported AuPd nanoparticles, prepared by the incipient wetness impregnation method, can effectively catalyze the oxidation of methane in water to produce methyl hydroperoxide, methanol, and  $\text{CO}_2$  using externally supplied  $\text{H}_2\text{O}_2$ . Moreover, the reaction can be performed using  $\text{H}_2\text{O}_2$  generated *in situ* from  $\text{H}_2$  and  $\text{O}_2$  introduced into the gas-phase reaction mixture. Notably, comparable productivity but significantly improved methanol selectivity was observed when employing  $\text{H}_2\text{O}_2$  generated *in situ*, relative to the use of preformed  $\text{H}_2\text{O}_2$ . This strategy not only simplifies the overall process by eliminating the need for external  $\text{H}_2\text{O}_2$  supply but also reduces operational costs and minimizes environmental impacts. The *in situ* generation of  $\text{H}_2\text{O}_2$  offers a more sustainable and economically viable oxidant alternative, while broadening potential applications in areas such as wastewater treatment, carbon capture and utilization, and the production of fine chemicals.

Tsubaki and co-worker<sup>135,136</sup> have reported an Au–Pd catalyst supported on carbon materials (CNTs, AC, and rGO) to convert methane into methane oxygenates with *in situ* generated  $\text{H}_2\text{O}_2$ . In particular, the CNT (carbon nanotube)-supported Pd–Au nanoparticles exhibit excellent catalytic performance in converting methane to methanol. They deduced that weak interaction resulting from CNT supports is more beneficial to the methanol productivity and selectivity. This strategy might facilitate the catalyst designs for this direct conversion of methane to methanol or other catalytic reactions in the future. They also discovered that the methanol productivity increases as the amount of Au loading in the Pd–Au/CNT catalyst increases. More in-depth investigation of this Pd–Au nanoparticle catalyst may facilitate the catalyst design and preparation for this direct



**Scheme 4** Current and proposed chemical processes for converting methane to methanol. Reproduced from ref. 127 with permission from the American Association for the Advancement of Science (AAAS), copyright 2017.



Table 3 Summary of various catalytic systems for coupling the oxidation of methane with *in situ* generation of H<sub>2</sub>O<sub>2</sub>

Ref.	Catalyst	Experimental conditions				Activity results								
		Catalyst mass (mg)	V <sub>solution</sub> (ml)	T (K)	Time (h)	Gas composition CH <sub>4</sub> /H <sub>2</sub> /O <sub>2</sub> /inert (bar)	Total pressure (bar)	Total product (mmol g <sub>metal</sub> <sup>-1</sup> h <sup>-1</sup> )	CH <sub>3</sub> OH amount (μmol)	CH <sub>3</sub> OH product (mmol g <sub>metal</sub> <sup>-1</sup> h <sup>-1</sup> )	Selectivity CH <sub>3</sub> OH	HCOOH	CH <sub>3</sub> OOH	CO <sub>2</sub>
132, 134	2.5%Au/2.5%Pd/TiO <sub>2</sub>	28	10	323.15	0.5	0.86/1.72/75.86/21.55	32	2.32	1.31	1.58	68.2%	0	15.1%	16.7%
135, 136	2.5%Au/2.5%Pd/CNTs	30	10	323.15	0.5	47/4/8/41	33	7.60	4.17	5.56	73.2%	22.3%	4.6%	0
142	AuPd@ZSM-5-C <sub>16</sub>	27	10	343.15	0.5	3.3/6.6/1.6/88.3	30	99.45	64.1	91.6	92.1%	4.3%	Trace	Trace
137	FeSO <sub>4</sub> + 1%Pd/C	0.37 mmol + 50	30	293.15	0.5	15/3/2.1/7.9	28	—	11	—	3.1%	60.6%	1.9%	34.4%
138	0.9%Fe-ZSM-5 + 1%Pd/AC	50 + 50	30	303.15	0.5	15/3/2.1/7.9	28	400 (Fe)	22	97.6	24.4%	57.8%	0	17.8%
139	0.9%Fe-ZSM-5 + 2%Pd/c-s-HCPP	50 + 25	30	323.15	1	15/3/2.1/7.9	28	185 (Fe)	24	51.8	28%	62%	0	11%

methane to methanol reaction or other catalytic processes in the future.

Park and coworkers<sup>137–139</sup> reported enhanced methane oxygenation when H<sub>2</sub> was introduced in the presence of both FeSO<sub>4</sub> and Pd/C. They attributed this to Fe<sup>2+</sup> accelerating hydroxyl radical ( $\cdot$ OH) formation, analogous to a Fenton-like mechanism. These radicals selectively oxidize methane to methanol. This highlights the superior efficiency of *in situ* H<sub>2</sub>O<sub>2</sub> generation over direct H<sub>2</sub>O<sub>2</sub> addition.

However, homogeneous Fe catalysts suffer from issues such as metal leaching and recovery challenges. To address this, Fe-ZSM-5 zeolites were explored as heterogeneous alternatives. While lower pH promoted H<sub>2</sub>O<sub>2</sub> formation and stability, ICP analysis revealed increased iron leaching under acidic conditions, compromising catalyst durability.<sup>140</sup>

To overcome these limitations, Park *et al.* developed a bifunctional system combining Fe-ZSM-5 with Pd/C supported on an acid-functionalized hyper-crosslinked porous polymer (Pd/c-s-HCPP). The acidic sites on HCPP facilitated *in situ* H<sub>2</sub>O<sub>2</sub> generation without external acid. This system achieved high methane conversion and selectivity under mild conditions, while enhancing catalyst stability.

Similarly, Yin *et al.*<sup>141</sup> designed Fe single-atom catalysts on N-doped carbon (Fe-SA/NC) with axial oxygen ligands (Fe–N<sub>4</sub>–O). *In situ* FTIR and DFT studies showed that Fe–O bonds polarized the C–H bond in methane, reducing activation energy to 0.8 eV. At 50 °C and 10 bar CH<sub>4</sub>/H<sub>2</sub>/O<sub>2</sub> (3:1:1), the catalyst delivered 98% methanol selectivity with a space-time yield of 1.2 g L<sup>-1</sup> h<sup>-1</sup>, outperforming enzymatic systems. The robust Fe–N coordination suppressed metal leaching, maintaining 95% activity after 100 cycles.

Furthermore, bimetallic Fe–Pd catalysts exhibit synergistic effects, enhancing the degradation of organic pollutants and recalcitrant contaminants. Fe facilitates electron transfer, while Pd accelerates key reaction steps, allowing efficient operation under mild conditions with reduced energy input. These systems not only remove toxic pollutants and heavy metals from wastewater but also enable resource recovery, promoting sustainable chemical processes.

Xiao and coworkers<sup>142</sup> recognized the importance of local H<sub>2</sub>O<sub>2</sub> concentration in methane oxidation by *in situ*-generated H<sub>2</sub>O<sub>2</sub> under mild conditions. Then, they proposed a molecular fence concept and designed a series of AuPd@zeolite-R catalysts. Au–Pd alloy nanoparticles were embedded in aluminosilicate zeolite crystals that rendered the external surface of the zeolite hydrophobic by appending organosilanes (R). Hydrogen, oxygen, and methane were easy to diffuse to the catalytically active sites to form H<sub>2</sub>O<sub>2</sub>, but the generated H<sub>2</sub>O<sub>2</sub> was difficult for diffusing out of the zeolite crystals. The hydrophobic sheath hinders H<sub>2</sub>O<sub>2</sub> diffusion from the encapsulated AuPd nanoparticles, enhancing its concentration in the zeolite crystals, thereby achieving excellent catalytic effects. The concept of a molecular fence opened a productive route to efficient catalysts for partial methane oxidation.



Several groups reported successful partial methane oxidation using  $\text{H}_2\text{O}_2$  generated *in situ* from hydrogen and oxygen and put forward some exciting concepts. However, the mechanism of partial methane oxidation should be further explored. We can design highly active catalysts that match the reaction with a complete understanding of the methane oxidation pathway.

## 6 Green adipic acid synthesis

Adipic acid, an important industrial dicarboxylic acid, has attracted considerable attention in recent years for its green synthesis, particularly *via* oxidation routes of cyclohexene or cyclohexane driven by *in situ*-generated  $\text{H}_2\text{O}_2$ . Sato *et al.*<sup>143</sup> pioneered a “green” route employing 30%  $\text{H}_2\text{O}_2$  for the direct oxidation of cyclohexene to high-purity adipic acid under mild conditions, achieving yields up to 93% while completely eliminating harmful  $\text{N}_2\text{O}$  emissions associated with traditional nitric acid methods. This work laid the foundation for  $\text{H}_2\text{O}_2$ -based adipic acid synthesis.

Ribeiro *et al.*<sup>144</sup> compared the oxidation performances of cyclohexene and cyclohexane, highlighting that the presence of the C=C double bond in cyclohexene facilitates adipic acid formation under  $\text{H}_2\text{O}_2$  and microwave-assisted conditions, reaching yields of 46%. In contrast, the activation of the C-H bond in cyclohexane is more challenging, resulting in lower efficiency. Their study also demonstrated partial recyclability of a C-chelated iron catalyst in ionic liquids, enhancing the sustainability of the catalytic system.

Van *et al.*<sup>145</sup> reviewed emerging catalytic processes for green adipic acid synthesis, emphasizing that coupling *in situ*  $\text{H}_2\text{O}_2$  generation with oxidation reactions improves efficiency and reduces risks associated with the storage and transportation of concentrated  $\text{H}_2\text{O}_2$ . Furthermore, Noyori *et al.*<sup>146</sup> summarized design principles and catalyst optimization strategies for aqueous-phase  $\text{H}_2\text{O}_2$  oxidation systems, advancing the development of tungstate and related catalysts.

From a biocatalytic perspective, Li *et al.*<sup>147</sup> reviewed peroxygenase-driven C-H functionalization reactions, focusing on *in situ*  $\text{H}_2\text{O}_2$  generation *via* electro- and photocatalytic methods. They highlighted the critical importance of precise control of  $\text{H}_2\text{O}_2$  concentration to enhance the reaction efficiency and prevent enzyme deactivation.

In summary, current research on green adipic acid synthesis mainly concentrates on designing efficient catalytic systems based on hydrogen peroxide and *in situ*  $\text{H}_2\text{O}_2$  generation technologies. By integrating traditional and biocatalytic approaches and optimizing reaction systems and catalysts, significant progress has been made toward environmentally friendly and sustainable industrial production of adipic acid.

## 7 Fenton process

When phenolic compounds are widely used as essential chemical raw materials and intermediates, a large amount of industrial phenol-containing organic wastewater that is difficult to degrade is generated. Untreated organic wastewater discharged directly into water bodies or soil can adversely affect aquatic organisms and soil diversity due to the presence of chemicals such as organic solvents, dyes, and pesticide residues. The Fenton synthesis process offers an economic and environmentally friendly solution for organic wastewater treatment through the efficient degradation and resource utilization of organic pollutants. It can convert organic components in wastewater into clean fuels, reducing environmental pollution, promoting circular economy development, and achieving dual benefits of wastewater treatment and resource utilization.

In recent years, there have been many studies on the Fenton reaction with  $\text{H}_2\text{O}_2$ , mainly Pd-based and Fe-based catalysts,<sup>148–157</sup> all of which can be used for the degradation of insoluble organic compounds and have achieved good results.

Fenton technology has made many advances in the field of water contaminant control. The current oxidant used,  $\text{H}_2\text{O}_2$ , is very costly, but its transportation and storage costs are incredibly high, so there is an urgent need to find a solution for it.

In analogy to reactions such as epoxidation of propylene and hydroxylation of benzene, perhaps the *in situ* participation of hydrogen and oxygen in the Fenton reaction compared to the direct participation of  $\text{H}_2\text{O}_2$ , the raw material of this system is not only easier to store but also allows the *in situ* production of  $\text{H}_2\text{O}_2$  from electrically generated  $\text{H}_2$  and  $\text{O}_2$  using catalysts such as Pd/C. These studies demonstrate the great potential of *in situ*-generated  $\text{H}_2\text{O}_2$  in waste treatment oxidation reactions, reducing the current costs associated with  $\text{H}_2\text{O}_2$  production. Furthermore, it is essential to highlight that the use of bimetallic Fe-Pd catalytic systems can improve the degradation performance and reduce the environmental impact of the process through the heterogeneity of Fenton ferrous ions and their associated synergistic effects.

## 8 Oxidative desulphurization

In numerous countries, coal has always been dominant, and the combustion of fossil fuels will bring severe environmental pollution, such as the issue of acid rain caused by  $\text{SO}_2$  emissions, which can easily disrupt the soil's acid-base balance, leading to a rapid reduction in crop yields and a severe threat to people's lives. Waste tires contain a large number of sulfur components. Many countries in Europe have banned tire disposal in landfills.<sup>158</sup> High sulfur contents reduce the quality of refined oil and poisonous catalysts, shortening their lifespan and activity. Additionally, sulfur pollution poses health risks, as prolonged exposure to



sulfur dioxide can cause respiratory diseases such as asthma and bronchitis. A notable example is the “Great Smog of London” in the mid-20th century, where extensive coal burning released large amounts of sulfur dioxide, leading to severe air pollution and thousands of deaths from respiratory and cardiovascular diseases. Therefore, reducing sulfur emissions is a critical issue. It is essential to promote energy conservation, implement strict sulfur dioxide emission controls, and develop cost-effective desulfurization technologies. Many researchers are dedicated to advancing desulfurization efforts.

The main application areas of oxidative desulfurization are addressing issues in the combustion of fossil fuel coal and pyrolysis of waste tires. Common desulfurization technologies encompass hydrodesulfurization, oxidative and biological desulfurization, adsorption, and ionic liquid extraction.<sup>159</sup> In the study of pyrolysis of waste tires, the central technology used in the past was hydrodesulfurization, which had disadvantages of high reaction temperature, high hydrogen pressure, large reactor volume, and high catalyst requirements.<sup>160</sup> Absolutely, oxidative desulfurization has emerged as a viable solution for mitigating sulfur-related issues in fuels. It utilizes H<sub>2</sub>O<sub>2</sub> as an oxidant, which is cost-effective and readily available. Notably, oxidative desulfurization offers environmental advantages over hydrodesulfurization; first, the technology is centered on selective oxidation, converting sulfides into soluble or separable sulfur derivatives, such as sulfuric acid or elemental sulfur, thereby achieving efficient desulfurization. It typically operates under relatively low-temperature and -pressure conditions, significantly reducing energy consumption. Second, oxidative desulfurization avoids common issues associated with absorption methods, such as the regeneration of absorbents and physical adsorption methods, such as adsorption saturation, resulting in relatively lower operating costs. Additionally, the byproducts of oxidative desulfurization are mostly non-toxic and recyclable substances, such as elemental sulfur, facilitating resource utilization. Compared with the traditional chemical desulfurization methods, it greatly reduces the risk of secondary pollution.<sup>161</sup> Indeed, Fe-based nanoparticles catalysts have shown promise in desulfurization processes.<sup>162</sup> Additionally, the inclusion of H<sub>2</sub>O<sub>2</sub> can enhance the desulfurization efficiency,<sup>160</sup> leading to shortened reaction times. H<sub>2</sub>O<sub>2</sub> exhibits the ability to effectively eliminate inorganic sulfur during coal combustion, contributing to cleaner and more efficient energy production.<sup>163</sup>

The utilization of supported catalysts has expanded to encompass the catalysis of thiophene compounds alongside H<sub>2</sub>O<sub>2</sub>. Studies have showcased remarkable desulfurization efficiency.<sup>158</sup> These catalysts exhibit recyclability across multiple reactions while maintaining stable activity. For instance, Li *et al.* used SiO<sub>2</sub>-supported H<sub>3</sub>PMo<sub>12</sub>O<sub>40</sub> and H<sub>2</sub>O<sub>2</sub> for catalyzing dibenzothiophene and diesel desulfurization, demonstrating sustained excellent performance even after

numerous cycles.<sup>164</sup> Xie *et al.*<sup>165,166</sup> enabled continuous oxidative desulfurization with 98.7% sulfur removal efficiency, leveraging *in situ* H<sub>2</sub>O<sub>2</sub> generation and instantaneous mixing at the microscale. Another study by Qi *et al.* focused on HPW/PEHA/Zr SBA-15 catalysis of dibenzothiophene, further contributing to the advancements in desulfurization processes.<sup>167</sup>

Absolutely, desulfurization technologies are rapidly evolving toward cost-effectiveness, reduced pollution, higher efficiency, and environmental friendliness. Presently, oxidative desulfurization technology remains a dominant method due to its effectiveness. However, other approaches such as adsorption, extraction, and biological desulfurization are also advancing, showing promising developments. This diversification in desulfurization techniques signifies a rapid expansion in the desulfurization industry.

## 9 Oxidation of sulfides to sulphones

Sulfone plays an essential role in modern organic chemistry. They are highly versatile components and can be used in various applications such as drugs, agrochemicals, or functional materials.<sup>168</sup> In recent years, sulfone electrolytes have gained great interest because of their wide electrochemical window and high safety.<sup>169</sup> Many natural organic synthesis products, drugs, and crop protection chemicals contain sulfone or sulfoxide functional groups. Sulfones and sulfoxides are widely used in pharmaceutical, agricultural, biological, chemical, and industrial fields because of their unique biological and chemical activities.

The four traditional and most common methods for the synthesis of sulfones are the oxidation of sulfides (or sulfoxides), the Foucault reaction with a sulfonyl chloride, the electrophilic capture of sulfoxides, and the addition of sulfonyl radicals to olefins or alkynes. Among them, the selective oxidation of sulfide is a direct method for preparing organic sulfoxide and sulfone.<sup>168</sup> Although a variety of other oxidants such as nitric oxide, different metal oxides, high valence iodine reagents, halides, or high halides, can be used to oxidize sulfides to sulfones, these preparation methods have issues with using toxic reagents, excessive reagent usage, low chemical selectivity or utilizing hazardous organic solvents. With people's attention to the environment, the search for environmentally friendly synthetic methods has been a research hotspot in recent years. In most studies, H<sub>2</sub>O<sub>2</sub> is commonly utilized as a “green” oxidant. The oxidation of sulfide to sulfone with H<sub>2</sub>O<sub>2</sub> as the oxide is mainly based on metal catalysts.

Trivedi<sup>170</sup> carried out simple and effective sulfonation oxidation with 30% H<sub>2</sub>O<sub>2</sub> as oxidant and VCl<sub>3</sub> as a catalyst. Studies have proved that tetrahydrofuran (THF) is a suitable solvent for selective oxidation of sulfide. When the solvent is THF, the sulfide yield is 83–98%.



Jain *et al.*<sup>171</sup> selected diphenyl sulfide as the substrate, 30% H<sub>2</sub>O<sub>2</sub> as the oxidant, and Nafion NR-50 as the catalyst to study its oxidation performance, optimize reaction conditions, and select solvent system. The results showed that dichloroethane was the most suitable solvent for the reaction when the catalyst loading was 0.25 g mmol<sup>-1</sup>. When the reaction time was 35 min, the yield was 98%. Jain *et al.* also studied the effect of catalyst loading by using different amounts of Nafion NR-50. It was found that when the catalyst loading was increased from 0.05 to 0.25 g mmol<sup>-1</sup>, the oxidation of diphenyl sulfide would significantly reduce the reaction time. Přeč and other coworkers<sup>172</sup> studied the oxidation of *p*-methyl phenyl sulfide (MPS) over layered titanium silicate catalysts with MFI and UTL-derived topology with H<sub>2</sub>O<sub>2</sub> as the oxidant. The results indicated that 95% selectivity of methyl phenyl sulfoxide with 40% conversion was achieved using the Ti-IPC-1-Pi catalyst. The selectivity for the oxidation of methyl phenyl sulfide to methyl phenyl sulfoxide is driven by diffusion restriction in the catalyst.

K. Bahrami *et al.*<sup>173</sup> studied the effects of catalyst dosage, H<sub>2</sub>O<sub>2</sub> dosage, solvent, and temperature on the catalytic oxidation of sulfide with 30% H<sub>2</sub>O<sub>2</sub> as the oxidant and BNPs-SiO<sub>2</sub>@(CH<sub>2</sub>)<sub>3</sub>NHSO<sub>3</sub>H as the catalyst. The results show that ethanol is the most suitable solvent for the reaction, and the reaction yield in ethanol solvent is 98%. When the molar ratio between sulfide and H<sub>2</sub>O<sub>2</sub> was 1/3, and 0.07 g BNPs-SiO<sub>2</sub>@(CH<sub>2</sub>)<sub>3</sub>NHSO<sub>3</sub>H reacted for 10 minutes, the yield was 98%.

## 10 Conclusions and future perspectives

The utilization of *in situ*-generated H<sub>2</sub>O<sub>2</sub> from H<sub>2</sub> and O<sub>2</sub> for selective oxidation presents a promising alternative to current industrial methods. This approach offers mild, eco-friendly reaction conditions, ensuring better control over the process while circumventing the by-products from high-temperature self-oxidation. Contrastingly, utilizing abundant molecular oxygen as a raw material eliminates the need to stockpile large quantities of H<sub>2</sub>O<sub>2</sub> on-site, effectively reducing production and equipment costs while sidestepping losses due to commercial H<sub>2</sub>O<sub>2</sub> decomposition. However, a key challenge arises from the inherently low yield of directly synthesized H<sub>2</sub>O<sub>2</sub> in the absence of stabilizers. The addition of stabilizers tends to influence the overall yield of H<sub>2</sub>O<sub>2</sub> while significantly impacting the longevity of catalysts and equipment. In particular, the introduction of stabilizers, such as acids and halogens, can compromise the operational lifespan of both the catalyst and the equipment. Overcoming these scientific hurdles is imperative for advancing synthesis reactions. Moreover, this method entails a dual-reaction process: H<sub>2</sub>O<sub>2</sub> synthesis and target substrate oxidation, involving a complex micro mechanism encompassing adsorption, desorption, and diffusion at the active site. Addressing H<sub>2</sub>O<sub>2</sub> decomposition is equally pivotal.

Therefore, future catalyst design should prioritize active sites capable of simultaneous dual reactions to minimize diffusion limitations on the generated H<sub>2</sub>O<sub>2</sub>. It has been proved that the integrated system generates many hydroperoxides and reactive oxygen species *in situ*, promoting the oxidation of inactive C–H bonds to corresponding oxygen-containing compounds. This observed enhancement surpasses the effects achieved by the direct addition of H<sub>2</sub>O<sub>2</sub>. It can be popularized and designed according to similar strategies, and can deepen the understanding of reaction mechanism, formulate different strategies, improve the overall product yield and realize the economic feasibility of the process route utilizing catalyst design, process parameters, and system control. In addition, enhancing the stability of the catalyst in the selective oxidation process, mitigating metal component loss or poisoning, considering the safety of the reaction system, and controlling the H<sub>2</sub>/O<sub>2</sub> mixture ratio in the system are the keys to achieve future industrial applications.

Moreover, electrocatalysis, photocatalysis, and similar methodologies are gaining prominence for their capability to generate H<sub>2</sub>O<sub>2</sub> *in situ* under milder conditions. These approaches hold promise as potentially milder and greener reaction pathways for the future. New catalyst designs, including perovskite-type catalysts and the integration of novel porous materials such as COFs, offer promising avenues for oxidation reactions coupled with *in situ* H<sub>2</sub>O<sub>2</sub> processes.

## 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.

## Author contributions

All authors have made substantial contributions to the conception, design, investigation, data analysis, or interpretation of this work and have approved the final version of the manuscript. Conceptualization: Jinghui Lyu, Xiaonian Li, and Chengrong Ding; methodology: Han Wu, Qingqing Li, and Shihao Wang; investigation: Jinke Yao, Tao Liu, Wenying Chu, and Feng Feng; data curation and analysis: Qunfeng Zhang, Qingtao Wang, and Dahao Jiang; writing – original draft preparation: Han Wu and Jinghui Lyu; writing – review and editing: Chengrong Ding, Xiaonian Li, and Guofu Zhang; funding acquisition: Xiaonian Li; supervision: Jinghui Lyu, Chengrong Ding, and Xiaonian Li; project administration: Chengrong Ding and Xiaonian Li.

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## Conflicts of interest

There are no conflicts to declare.



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