

Review

Recent Advances of Oxygen Carriers for Hydrogen Production via Chemical Looping Water-Splitting

Wenxi Chang ^{1,2}, Yue Hu ^{2,3}, Weibin Xu ^{2,3}, Chuande Huang ^{2,*} , Haonan Chen ¹, Jiahui He ¹, Yujia Han ^{2,3}, Yanyan Zhu ^{1,*}, Xiaoxun Ma ¹ and Xiaodong Wang ^{2,*}

- ¹ School of Chemical Engineering, Northwest University, International Scientific and Technological Cooperation Base of MOST for Clean Utilization of Hydrocarbon Resources, Chemical Engineering Research Center for the Ministry of Education for Advance Use Technology of Shanbei Energy, Xi'an 710069, China
- ² CAS Key Laboratory of Science and Technology on Applied Catalysis, Dalian Institute of Chemical Physics, Chinese Academy of Sciences, Dalian 116023, China
- ³ School of Chemical Engineering, University of Chinese Academy of Sciences, Beijing 100049, China
- * Correspondence: huangchuande@dicp.ac.cn (C.H.); zhuyanyan@nwu.edu.cn (Y.Z.); xdwang@dicp.ac.cn (X.W.)

Abstract: Hydrogen is an important green energy source and chemical raw material for various industrial processes. At present, the major technique of hydrogen production is steam methane reforming (SMR), which suffers from high energy penalties and enormous CO₂ emissions. As an alternative, chemical looping water-splitting (CLWS) technology represents an energy-efficient and environmentally friendly method for hydrogen production. The key to CLWS lies in the selection of suitable oxygen carriers (OCs) that hold outstanding sintering resistance, structural reversibility, and capability to release lattice oxygen and deoxygenate the steam for hydrogen generation. Described herein are the recent advances in designing OCs, including simple metal oxides (e.g., Fe, Zn, Ce, and Ti-based metal oxides) and composite metal oxides (e.g., perovskite, spinel, and garnets), for different CLWS processes with emphasis on the crucial parameters that determine their redox performance and future challenges.

Keywords: chemical looping; water-splitting; hydrogen; oxygen carrier; redox performance



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1. Introduction

An urgent desire to reduce reliance on fossil fuels has been spurred in recent years due to the steeply increased concentration of greenhouse gases in the atmosphere [1]. Hydrogen, as an alternative, has been regarded as a promising fuel that possesses a high calorific value (121.00 MJ/kg) that is 2.8 times and four times greater than petroleum and coal, respectively, while giving clean water as the only combustion product [2–4]. More importantly, hydrogen is an important chemical raw material for various industrial processes, such as crude oil refining, synthesis of ammonia and methanol, etc. [5]. Currently, the production of hydrogen is mainly based on fossil fuels, such as steam methane reforming (SMR) and coal gasification (CG) [2,6,7]. However, both commercial routes are energy-intensive, rendering massive CO₂ emissions reaching as high as 10–16 and 22–35 kgCO₂e/kgH₂ for the SMR and CG processes, respectively [8–11]. Although carbon capture and storage technologies can potentially reduce the CO₂ emissions, this would inevitably induce a significant increase in energy penalty and manufacturing costs [12,13].

As a reverse reaction of hydrogen combustion, hydrogen production directly from water-splitting is considered a sustainable approach. Intuitively, direct thermal splitting of water (H₂O(l) → H₂(g) + 1/2O₂(g), ΔH⁰₂₉₈ = 286 kJ/mol) is the simplest method to implement this goal. However, thermodynamic analysis shows that a positive Gibbs free energy is maintained for this reaction until a temperature higher than 4370 K (Figure 1), revealing an extremely tough process to break the O–H bond in H₂O molecules [14].

Kogan et al. [15] proposed that water conversion only reached ca. 25% at a temperature of 2500 K and pressure of 0.05 bar, and rapid cooling treatment was needed to suppress the reverse reaction between generated H_2 and O_2 , causing an increased energy burden. A study showed that hydrogen produced by pyrolyzing the mixture of water and argon was lower than 3% percentage of all off-gases at 2000–2500 °C [16]. Recent reports suggested that water-splitting driven by light and electricity enabled production of hydrogen at ambient temperature [17,18]. Compared with the photocatalytic process that is still in its infancy, electrochemical water-splitting could provide roughly 5% of the world's hydrogen energy [19]. The purity of hydrogen generated in the electrolysis process is high, but the electrolysis process consumes a large amount of power, and the electricity cost accounts for around 80% of the overall water electrolysis process, rendering the scale-up of this technology strongly depending on the availability of low-cost and green electricity [20–22]. Therefore, research is ongoing to explore more efficient methods for hydrogen production with low carbon emissions and energy penalty.

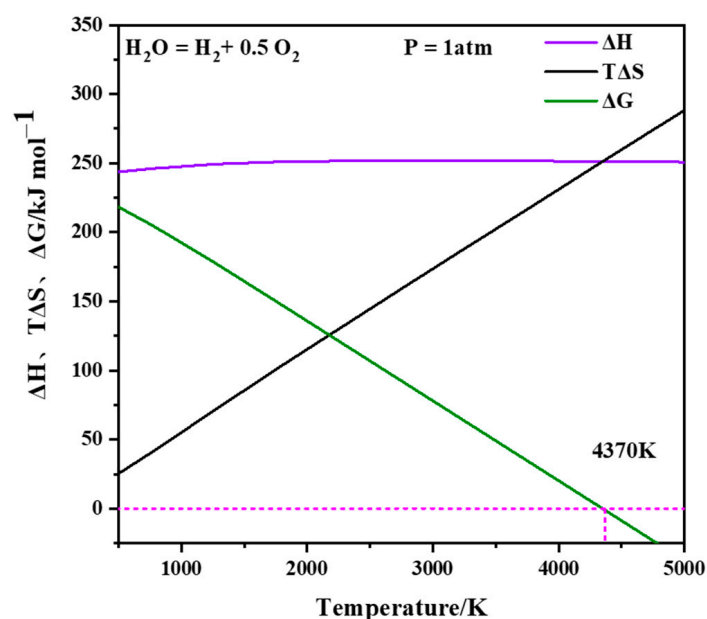


Figure 1. Variation of ΔH , $T\Delta S$, and ΔG with the temperature for direct water-splitting at 1 atm. Adapted with permission from Ref. [14], copyright 2007, American Chemical Society.

Chemical looping technology that was initially designed as an advanced combustion method has shown great potential for energy storage and production of chemicals due to the virtue of process intensification, leading to improved process efficiency and notably reduced CO_2 emission and investment expenses [23–25]. In recent years, pioneering reports have verified that applying the chemical looping concept to water-splitting, known as chemical looping water-splitting (CLWS), enables efficient hydrogen production at temperatures below 1300 °C. Compared with continuous reaction mode, CLWS reaction, utilizing metal oxides as oxygen carriers (OCs), is fulfilled by two or three spatiotemporal separated steps, including abstracting lattice oxygen from OCs via reduction treatment, regeneration by water with production of hydrogen, and in some cases, further oxidation by air to fully recover the OCs. Such a multistep reaction not only affords great convenience in separating hydrogen from products, but also requires less useful work and reaction free energy changes for water decomposition [26,27]. Furthermore, the introduction of OCs as redox catalysts significantly lowers the threshold to break the O–H bond of H_2O and improves the reaction kinetics [28–30].

In general, the yield of hydrogen in a single redox cycle is positively correlated with the oxygen capacity of OCs, which in turn is closely related to the reduction method to remove the lattice oxygen. According to the distinct strategies in abstracting lattice oxygen

from OCs, CLWS can be classified into processes including a two-step thermochemical water-splitting cycle (thermal reduction, Figure 2A) [31], a methane chemical looping process (CH_4 reduction, Figure 2B), a chemical looping water gas shift reaction (CO reduction, Figure 2C) [32], a syngas (mixture of H_2 and CO) chemical looping process (syngas reduction, Figure 2D) [33], and a photo-thermochemical cycle (photochemical reduction, Figure 2E) [34]. As for CLWS reaction, the economic efficiency of hydrogen production is substantially determined by redox properties of OCs, including mechanical strength, oxygen capacity, reactivity, selectivity, and structural stability, as well as low toxicity and low cost [35,36]. A series of reviews have been proposed in recent years that comprehensively summarized the potential applications of chemical looping technology [37], reaction mechanism of OCs [38], and the recent advantages in OCs for chemical looping conversion of alkanes [39–41]. As for hydrogen production via the chemical looping method, Voitic et al. [42] have reviewed the progress with emphasis on the iron-based OCs, while Luo et al. [43] detailed the system integration, economic analysis, and OCs for the chemical looping reforming process (cofeeding the fuel and water to reactor) and fuel-driven chemical looping water-splitting. Herein, the review gives a detailed survey on the current state of OC development for hydrogen production via different CLWS processes. The effect of the inherent properties of OCs on feed-gas conversion or product selectivity and future challenges in designing the OCs are summarized.

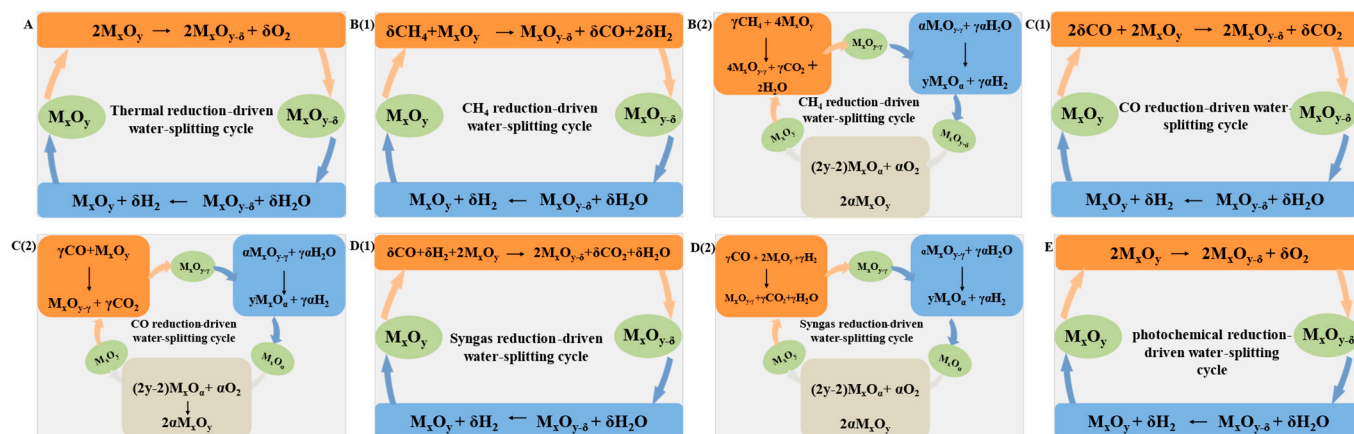


Figure 2. Chemical looping water-splitting process with various reducing methods: (A) two-step water-splitting thermochemical cycle, (B) methane chemical looping process, (C) chemical looping water gas shift process, (D) syngas chemical looping process, and (E) photo-thermochemical cycle.

2. Processes for Chemical Looping Water-Splitting

2.1. Two-Step Thermochemical Water-Splitting

To address the problems of high reaction temperature and low conversion for direct water-splitting, in 1966 Funk et al. [26] proposed the strategy of two-step thermochemical water-splitting (TCWS) for hydrogen production. The key to this process lies in selecting suitable metal oxides as OCs, which are thermally reduced at high temperature ($>1300\text{ }^{\circ}\text{C}$) to release oxygen and subsequently recovered by H_2O at a temperature generally lower than $1100\text{ }^{\circ}\text{C}$ with production of hydrogen. As a result, direct contact between hydrogen and oxygen is avoided, which bypasses the issue of hydrogen separation and enables higher water conversion at lower temperatures. In particular, when concentrated solar energy is utilized to heat the OCs (two-step solar thermochemical cycles), the solar energy is transformed to chemical energy with a theoretical solar-to-fuel efficiency reaching higher than 70% [14]. However, the high reaction temperature and thermal shocks induced by temperature swing pose a huge challenge in designing the OCs [44,45].

The $\text{Fe}_3\text{O}_4/\text{FeO}$ redox pair was one of the earliest OCs used in a two-step TCWS cycle. Based on the thermodynamic analyses, Nakamura et al. [46] and Steinfeld et al. [47] reported that it was feasible to apply $\text{Fe}_3\text{O}_4/\text{FeO}$ for hydrogen generation via thermochemical cycle

at temperatures of ca. 2000 °C (Figure 3). However, these oxides were subjected to severe sintering under high operating temperatures. To address this issue, some active metals or inert supports were introduced to the OCs in an attempt to lower the thermal reduction temperature or improve their sintering resistance. Han et al. [48] found that removal of lattice oxygen from ferrites was realized at lower temperature of 1200 °C after doping of Ni, Mn, and Cu while water-splitting was implemented at 800 °C. Among the screened materials, NiFe_2O_4 displayed the highest rate for hydrogen production with an average H_2 yield of 0.442 mL/g (the amount of produced H_2 per gram of oxygen carrier in each cycle). Structural analysis revealed that no obvious changes occurred after these redox reactions, which demonstrated a high stability of this material. Kodama et al. [49,50] also suggested that Ni-modified ferrites in $\text{A}_x\text{Fe}_{3-x}\text{O}_4$ ($\text{A} = \text{Mn}, \text{Co}, \text{Ni}$) exhibited better reactivity than Co or Mn promoters, and a scaling relationship between Ni content (from 0 to 1) with corresponding performance was noticed. When Ni loading increased to $x = 1$, a maximum hydrogen productivity of ca. 10 mL/g is reached. Their follow-up studies showed that loading of ferrites on ZrO_2 support significantly slowed down the sintering process, leading to stable hydrogen production. Ishihara et al. [51] reported that combination of NiFe_2O_4 with YSZ (yttrium-stabilized zirconia) support was capable of promoting the redox stability and oxygen mobility, resulting in a negligible decline in hydrogen production in 10 continuous cycles with reduction temperature of 1773 K and water-splitting temperature of 1473 K. Miller et al. [52] proposed that a mass ratio of 1/3 for $\text{Co}_{0.67}\text{Fe}_{2.33}\text{O}_4/\text{YSZ}$ could effectively inhibit the agglomeration of OCs and enable stable hydrogen production during eight cycles. In contrast, the bare $\text{Co}_{0.67}\text{Fe}_{2.33}\text{O}_4$ oxides was inactivated in the second cycle.

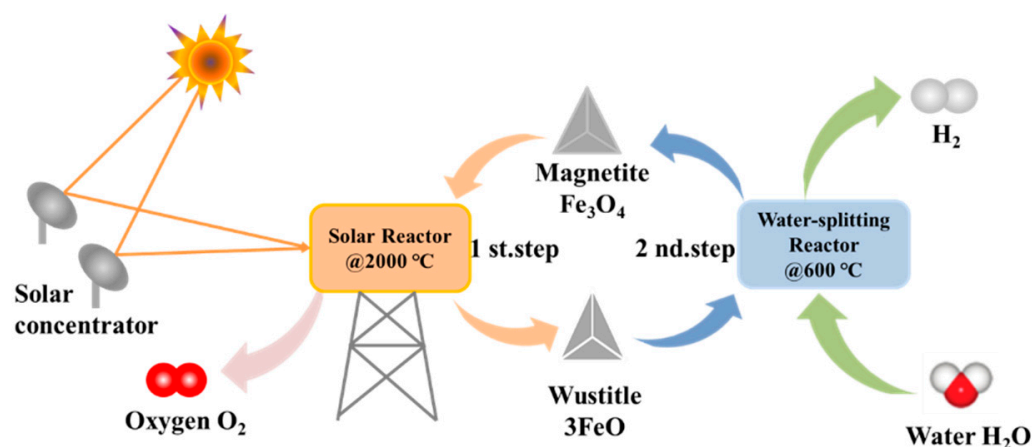


Figure 3. Schematic for two-step solar thermochemical water-splitting using $\text{Fe}_3\text{O}_4/\text{FeO}$ redox system. Adapted with permission from Ref. [47], Copyright 1999, Elsevier.

In 1977, Bilgen's team [16] applied a ZnO/Zn redox pair to the two-step TCWS cycle. Experimental results proved that thermal decomposition of ZnO into gaseous Zn (boiling point of 907 °C) and O_2 was achieved at 2338 K using a solar furnace. Thermodynamic calculations showed that the theoretical solar-to-fuel efficiency (without heat recovery) of the water-splitting cycle could reach 51% when the redox cycle was conducted with thermal reduction temperature of 2338 K (ZnO decomposition) and water-splitting at 1500 K. However, practical application of ZnO in TCWS remains greatly challenging due to the facile recombination of Zn and O_2 before introduction of water; thus, a quench treatment is necessary to separate Zn from O_2 by rapid condensation of Zn gas into solid form. The numerical simulation results by Palumbo et al. [53] showed that a conversion of 80% was achieved for ZnO decomposition when a high solar flux density of $2 \text{ MW}/\text{m}^2$ and quenching rate of $2.0 \times 10^7 \text{ K/s}$ were applied. Another challenge for application of ZnO/Zn redox pairs lies in the formation of a ZnO shell during the water-splitting reaction ($\text{Zn} + \text{H}_2\text{O} \rightarrow \text{ZnO} + \text{H}_2$), which significantly slows down the reaction kinetics [54].

Therefore, recent works have been devoted to kinetic study on this process with the aim to improve the hydrogen production efficiency. Ernst et al. [55] investigated the reaction between submicron zinc and ~50 mol% H₂O/Ar at 603–633 K. It was found that the initial reaction was a surface-controlled process with a fast rate and apparent activation energy of 42.8 ± 7.4 kJ/mol, while the subsequent reaction was a slow diffusion control process due to the restriction of surface ZnO layer on the migration of Zn²⁺. Lv et al. [56] investigated the effect of particle size on the water-splitting process and found that zinc powder with different sizes exhibited similar hydrolysis mechanisms, but that reducing the particle size could significantly enhance the reaction rate.

CeO₂, as a benchmark OC for two-step TCWS, has attracted particular attention since 2006 due to its outstanding redox properties. Abanades et al. [57] found that Ce₂O₃ obtained by thermal reduction of CeO₂ can be facilely oxidized by water with a conversion reaching 100%. However, a temperature of above 2000 °C is needed for complete reduction of CeO₂ to Ce₂O₃, which inevitably resulted in serious sintering of OCs. To bypass this issue, in most cases for TCWS applications, the reduction temperature was lowered to below 2000 °C, leading to formation of non-stoichiometric CeO_{2-δ}. Steinfeld et al. [58] reported that 2.5–2.9 mL of O₂ can be released per gram of ceria in a single redox cycle (2.5–2.9 mL/g), corresponding to δ value of 0.039–0.044, when thermal reduction was conducted at ca. 1600 °C. This rendered a hydrogen evolution rate reaching a peak value of 7.6×10^2 mL/min during the following water-splitting step (<900 °C), which resulted in an overall solar-to-hydrogen efficiency of 0.7%. Costa Oliveira et al. [59] investigated the effect of CeO₂ morphology on water-splitting reaction and discovered that H₂ yield over three-dimensionally ordered macroporous (3DOM) CeO₂ was approximately 32% greater than that of conventional CeO₂ foam, suggesting that engineering the morphology was effective in adjusting the reaction kinetics and oxygen mobility. Although much progress has been achieved these years, the low oxygen capacity and high reduction temperature of pure ceria greatly limited the hydrogen productivity and process efficiency, which impeded the feasibility of ceria in practical STWS applications. To address this issue, significant research has been performed to tailor the redox properties of Ce-based OCs by tuning the oxygen vacancies and lattice strain via doping foreign cations [60–63]. Gokon et al. [64] compared the performance of 30 mol% M-doped CeO₂ (M = Fe, Co, Ni, or Mn) in two-step TCWS cycles and found that all dopants except for Ni greatly increased the oxygen evolution during thermal reduction at 1500 °C. However, the yield of H₂ is lower than that of bare CeO₂, which may be ascribed to severe sintering of these OCs at high temperature. Tamaura et al. [65] investigated the performance of CeO₂-MO_x solid solution (M = Mn, Fe, Ni, Cu) with Ce/M molar ratio of 9 for two-step TCWS. As shown in Figure 4A, introducing of di- or tri-valent transition metal cations into CeO₂ notably promoted H₂ generation by enhancing releasing of lattice oxygen. The same group further screened a series of Ce_{0.9}M_{0.1}O_{2-δ} ceramics (M = Mg, Ca, Sr, Sc, Y, Dy, Zr, and Hf), and found that the evolution of O₂ (reduction of Ce⁴⁺ to Ce³⁺) in the thermal reduction step was improved by doping cations with higher valence state and smaller effective ionic radius (Figure 4B) [66,67]. Furthermore, the amount of H₂ produced in the water-splitting step was dominated by the amount of O₂ released in the reduction step. Instead, Abanades and coworkers [68] showed that addition of tri-valent dopants, including Y, La, Pr, and Gd, mainly contributed to improve the sintering resistance rather than the reducibility of ceria, but the ceria/zirconia mixed oxide was capable of greatly enhancing the reduction of bulk Ce⁴⁺. After detailed study on the correlation between zirconium content and redox properties of Ce_{1-δ}Zr_δO₂ (0 ≤ δ ≤ 0.5), it was found that the ratio of Ce³⁺/Ce⁴⁺ in thermally reduced oxides gradually increased with Zr content while an optimal composition of Ce_{0.75}Zr_{0.25}O₂ gave the highest hydrogen yield of 6.67 mL/g [69,70]. Their follow-up studies suggested that decoration of CeO₂-ZrO₂ with 1% Gd results in further promotion of hydrogen production to 7.58 mL/g [71]. Instead of exploring the nonstoichiometric chemistry that results in partial reduction of Ce⁴⁺ to Ce³⁺, Wang et al. [72] discovered that Sn-modified oxide of CeO₂-xSnO₂ (x = 0.05–0.20) was converted to thermodynamically

cally stable $\text{Ce}_2\text{Sn}_2\text{O}_7$ during the thermal reduction step (1400°C), which enabled deep reduction of Ce^{4+} to Ce^{3+} rather than $\text{CeO}_{2-\delta}$. When the Sn content was optimized to $x = 0.15$, a hydrogen production as high as 7.18 mL/g (Figure 5), 3.8 times greater than that of unmodified CeO_2 , was obtained.

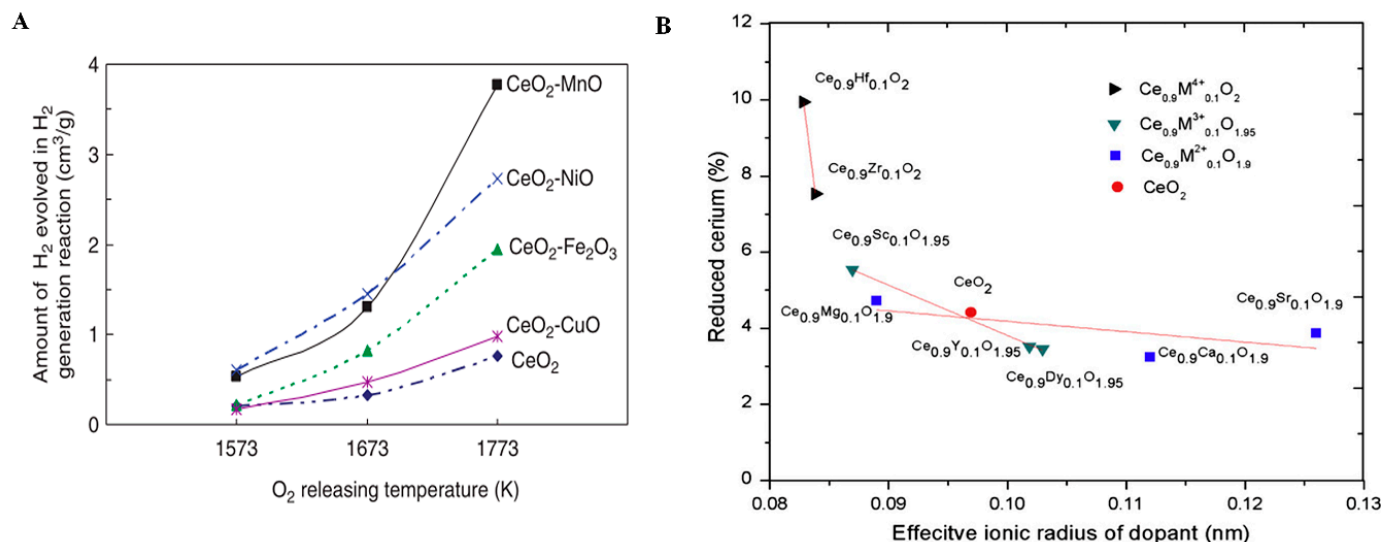


Figure 4. (A) Correlation between O_2 -releasing temperature and the amount of hydrogen produced in the reaction with $\text{CeO}_2\text{-MO}_x$. Reprinted with permission from Ref. [65], copyright 2007, Elsevier. (B) Effect of effective ionic radius and valence state of dopants on the reduction degree of cerium. Reprinted with permission from Ref. [66], copyright 2011, Elsevier.

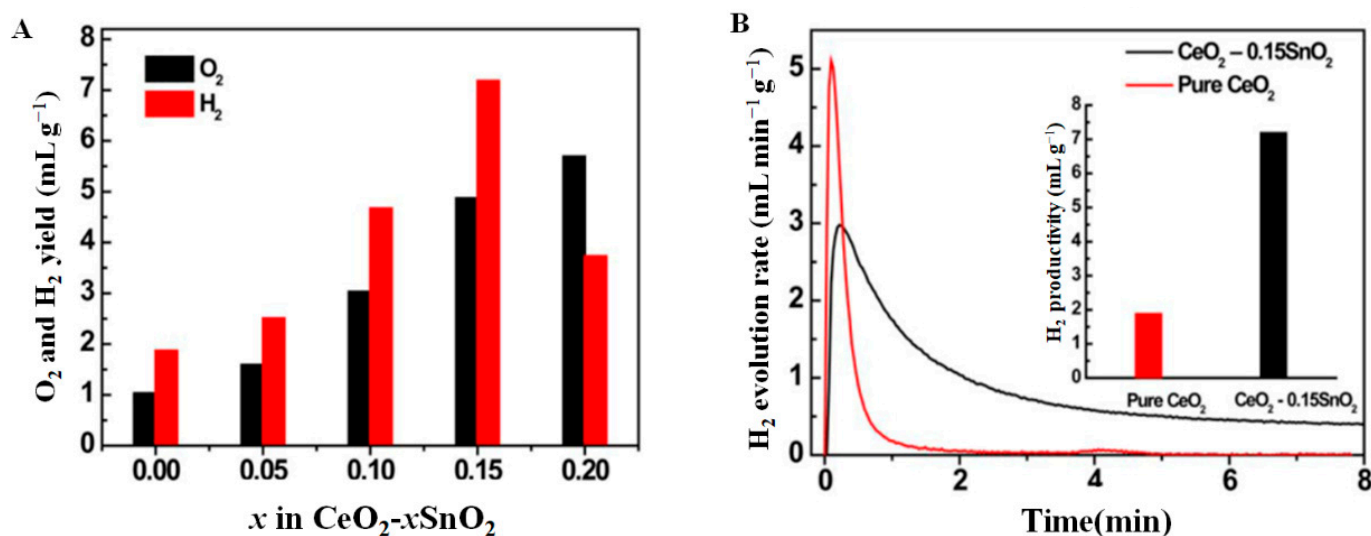


Figure 5. (A) Evolution of O_2 and H_2 for $\text{CeO}_2\text{-}x\text{SnO}_2$ ($x = 0.05\text{--}0.20$) during redox reactions. (B) H_2 evolution rate during water-splitting reaction at 800°C for $\text{CeO}_2\text{-}0.15\text{SnO}_2$ and pure CeO_2 . Adapted with permission from Ref. [72], Copyright 2017, Wiley.

Perovskite oxides (ABO_3) represent an important class of composite materials, wherein the A-site is generally occupied by rare earth, alkaline earth, or alkali metals that coordinate with 12 oxygen anions while the B-site is typically occupied by transition metals located in the octahedral interstices of the oxygen framework. These oxides hold virtue of outstanding redox properties, structure reversibility, and high-temperature stability, which render these oxides promising OCs for two-step TCWS reactions [73,74]. McDaniel et al. [75] discovered that Sr/Al doped LaMnO_3 exhibited exceptional oxygen mobility, which released 8 times more O_2 than CeO_2 at 1350°C , while the temperature for lattice oxygen

desorption was nearly 300 °C lower than that of CeO₂. Consequently, a hydrogen yield 9 times greater than that of CeO₂ is obtained at 1000 °C. R. Barcellos and colleagues [76] demonstrated that doping of Ce at B-site of BaMnO_{3-δ} allowed optimization of the oxygen vacancy formation energy. Among the investigated BaCe_xMn_{1-x}O_{3-δ} samples (0 ≤ x ≤ 1), BaCe_{0.25}Mn_{0.75}O_{3-δ} displayed the best water-splitting performance and yielded nearly 3 times more hydrogen than ceria when reduced at 1350 °C. Similarly, Li et al. [31] also found that introduction of Co into LaGaO₃ notably reduced the oxygen vacancy formation energy to ca. 3.0 eV, which was favorable for both lattice oxygen desorption and subsequent water-splitting. The best LaGa_{0.4}Co_{0.6}O_{3-δ} sample exhibited a H₂ yield of 0.478 mmol/g that was 15 times greater than CeO₂ (0.032 mmol/g) when reduced at 1350 °C and re-oxidized at 800 °C. Carter et al. [77] screened a series of Ca_{0.5}Ce_{0.5}MO₃ oxides (M = Sc, Ti, V, Cr, Mn, Fe, Co, and Ni) for a two-step solar thermochemical process using density functional theory (DFT) calculations. The calculation results suggested that the oxygen vacancy formation energy of Ca_{0.5}Ce_{0.5}MnO₃ and Ca_{0.5}Ce_{0.5}FeO₃ was slightly lower than that of CeO₂ (4.0~4.3 eV) and located in the range of 3.65~3.96 and 3.77~4.06 eV, respectively, which fitted well to the target window (3.2~4.1 eV) for thermochemical cycles (Figure 6). Qian et al. [78] found that the CaTi_{0.5}Mn_{0.5}O_{3-δ} was favorable for oxygen releasing and water-splitting due to its intermediate reduction enthalpy value of 200–280 KJ/mol-O (the standard enthalpy to remove one mole lattice oxygen of OC) and large entropy value of 120–180 J/mol-O·K (the standard entropy to remove one mole lattice oxygen of OC), which enables a hydrogen production of 10.0 mL/g in redox cycle between 1350 °C (thermal reduction) and 1150 °C (water-splitting) with a short cycle time of 1.5 h. All these results indicate that perovskite oxides are among the best candidates for two-step TCWS processes, and that A/B site engineering is efficient in modulating their redox properties. However, the step forward for practical application of this technology is still greatly hindered by high operating temperature and relatively low hydrogen yield due to the low oxygen redox capacity and long temperature swing time, which should be resolved ultimately by developing more effective OCs.

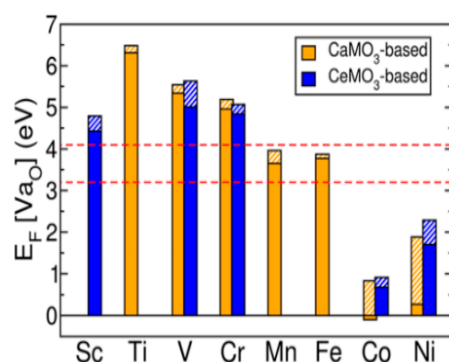


Figure 6. Oxygen vacancy formation energy of Ca_{0.5}Ce_{0.5}MO₃ perovskites derived from CaMO₃ (orange bars) and CeMO₃ (blue bars) oxides. Red dashed lines show the target scope for solar thermochemical hydrogen production (3.2~4.1 eV). Reprinted with permission from Ref. [77], copyright 2020, American Chemical Society.

2.2. Methane Chemical Looping Process

The two-step TCWS cycle approach has substantially decreased the reaction temperature (≤1500 °C) compared to direct thermal splitting of water, but the high reaction temperature still puts forward tough requirements for the reaction equipment. Furthermore, the continuous temperature swing and thermal shock during the cyclic reaction not only reduces the reaction efficiency but also leads to rapid decrease of redox performance, which seriously limits the commercialization of this technology. To this end, recent reports verified that introduction of reducing gas could significantly accelerate the reduction kinetics of OCs, lower the reaction temperature to below 1000 °C, and increase the amount of lattice oxygen desorption, substantially enhancing the hydrogen production efficiency [79–83].

Methane, the main component of natural gas, has been widely studied due to its abundant reserves, high hydrogen-to-carbon ratio, and strong reducibility [84,85]. The two-step TCWS cycle with methane as reducing gas is commonly referred to as methane chemical looping process (Figure 2B). The facile reduction of OCs in methane atmosphere enables a closed reaction loop at much milder conditions, which potentially reduces the energy penalty and improves the economics of this process. Compared with conventional standalone systems that generate hydrogen and electricity separately based on methane, it is assessed that chemical looping hydrogen generation with methane as reduction gas was able to save more than 16% of energy input while reducing beyond 98% of CO₂ emissions, rendering a low cost of \$32.87/MWh for H₂ production [86].

The methane conversion over OCs undergoes different processes, including partial oxidation or total combustion, when modulating the redox properties of these oxides. As for OCs with relatively low reducibility, syngas production is favored by selective methane oxidation ($\text{CH}_4 + \text{O}_L \rightarrow \text{CO} + 2 \text{H}_2$) (Figure 2B1). The reduced OCs can be subsequently recovered by water oxidation with generation of hydrogen ($\text{H}_2\text{O} \rightarrow \text{O}_L + \text{H}_2$). The overall reaction is generally referred as chemical looping steam methane reforming (CL-SMR) [87]. Compared with traditional steam methane reforming reaction ($\text{CH}_4 + \text{H}_2\text{O} \rightarrow \text{CO} + 3 \text{H}_2$), the CL-SMR process can obtain syngas with H₂/CO ratio of 2, which is suitable for Fischer-Tropsch synthesis and methanol. Furthermore, pure hydrogen is produced separately from syngas, bypassing the gas separation process; more importantly, through the rapid cycling of these OCs in different reactors, coking deposition on the catalyst is greatly inhibited [88–91]. When OCs with high reducibility are applied, methane is fully oxidized to CO₂ and H₂O ($\text{CH}_4 + 4 \text{O}_L \rightarrow \text{CO}_2 + 2 \text{H}_2\text{O}$), which is a strong exothermic reaction, and the released heat can be used for the next step water-splitting reaction, achieving heat self-sufficiency. Another dramatic difference between such a process with CL-SMR is that a third step by air oxidation is normally required to fully regenerate the OCs due to their high oxidation potential (Figure 2B2). This process (three reactors chemical looping M-TRCL) enables hydrogen production with low CO₂ emissions, since the high concentration of CO₂ can be captured by simple condensation [92].

2.2.1. Supported or Doped Iron Oxides

As for methane-driven three step CLWS, the studied OCs are mainly focused on iron oxides, which holds the virtue of low cost and environmentally friendly features [93,94]. However, developing an efficient method to avoid coke formation, which can degrade the hydrogen purity in the water-splitting step, over these OCs during methane atmosphere remains a great challenge [95]. Ku et al. [96] reported that methane dissociation and coke formation occurred over Fe₂O₃/Al₂O₃ OCs due to the generation of the FeAl₂O₄ phase, since the newly formed phase displayed poor oxygen mobility and rendered notably decreased oxygen capacity of OC for chemical looping reactions. To address this issue, Xiang et al. [97] proposed that loading Fe₂O₃ to MgAl₂O₄ support greatly suppressed the solid-phase reaction between iron oxides and the support, thereby restraining the carbon formation from CH₄ dissociation. Furthermore, they found that K-promoted Fe₂O₃/Al₂O₃ further improved the resistance towards coke deposition by decreasing the reduction activity of oxygen carrier. Li et al. [98] found that a combination of mixed ionic-electronic conductive (MIEC) support of La_{0.8}Sr_{0.2}FeO₃ with Fe₂O₃ improved lattice oxygen diffusion from the bulk to surface, which promoted the elimination of carbon during the methane reduction process.

Instead, when syngas is set as aimed products (methane-driven two-step CLWS), the proper OC needs to bear sufficient reactivity for selective methane conversion. There is usually a seesaw effect in reactivity and selectivity due to the proportional relationship between C–H bond activation ability of CH₄ and oxidation ability [99]. For instance, common iron oxides (Fe₂O₃ and Fe₃O₄) are reactive towards methane conversion, but CO₂ production is preferred over these OCs due to the high oxygen capacity [100,101]. To improve the selectivity of syngas at the CL-SMR process, extensive efforts have been

made to break the seesaw effect. Fan et al. [102] proposed that decreasing the particle size of Fe_2O_3 to 3–5 nm by confining these oxides in mesoporous SBA-15 ($\text{Fe}_2\text{O}_3/\text{SBA-15}$) enabled notable enhanced syngas selectivity to above 99%. Corresponding DFT calculations demonstrated that the low coordinated Fe and O atoms of Fe_2O_3 nanoparticles promoted CH_4 activation, cleavage of Fe–O bond, and formation of CO while suppressed the further oxidation to CO_2 (Figure 7). Based on DFT + U calculations, they further analyzed the mechanism for CH_4 partial oxidation over $\alpha\text{-Fe}_2\text{O}_3$ and claimed that increasing the oxygen vacancy concentration decreased the activation barriers for breaking the C–H bonds of CH_4 and Fe–C bonds, which contributed to improved reactivity and syngas selectivity (Figure 8) [103]. Recently, Wang’s team [104] discovered that Y-modified $\text{Fe}_2\text{O}_3/\text{Al}_2\text{O}_3$ (Y: Fe = 1.5) could produce a new phase of $\text{Y}_3\text{Fe}_2\text{Al}_3\text{O}_{12}$ garnet after high temperature treatment (1200°C), which increased the dispersion of active metal Fe, and changed the coordination environment of oxygen species. Lattice oxygen could coordinate with Fe, and also coordinate with Al and Y (Fe–O–Al, Fe–O–Y), which renders increased oxygen vacancy formation energy from 1.4 eV for bare Fe_2O_3 to 1.997 eV of $\text{Y}_3\text{Fe}_2\text{Al}_3\text{O}_{12}$, which substantially avoids excessive oxidation of CH_4 to CO_2 while maintaining CO selectivity as high as 98%.

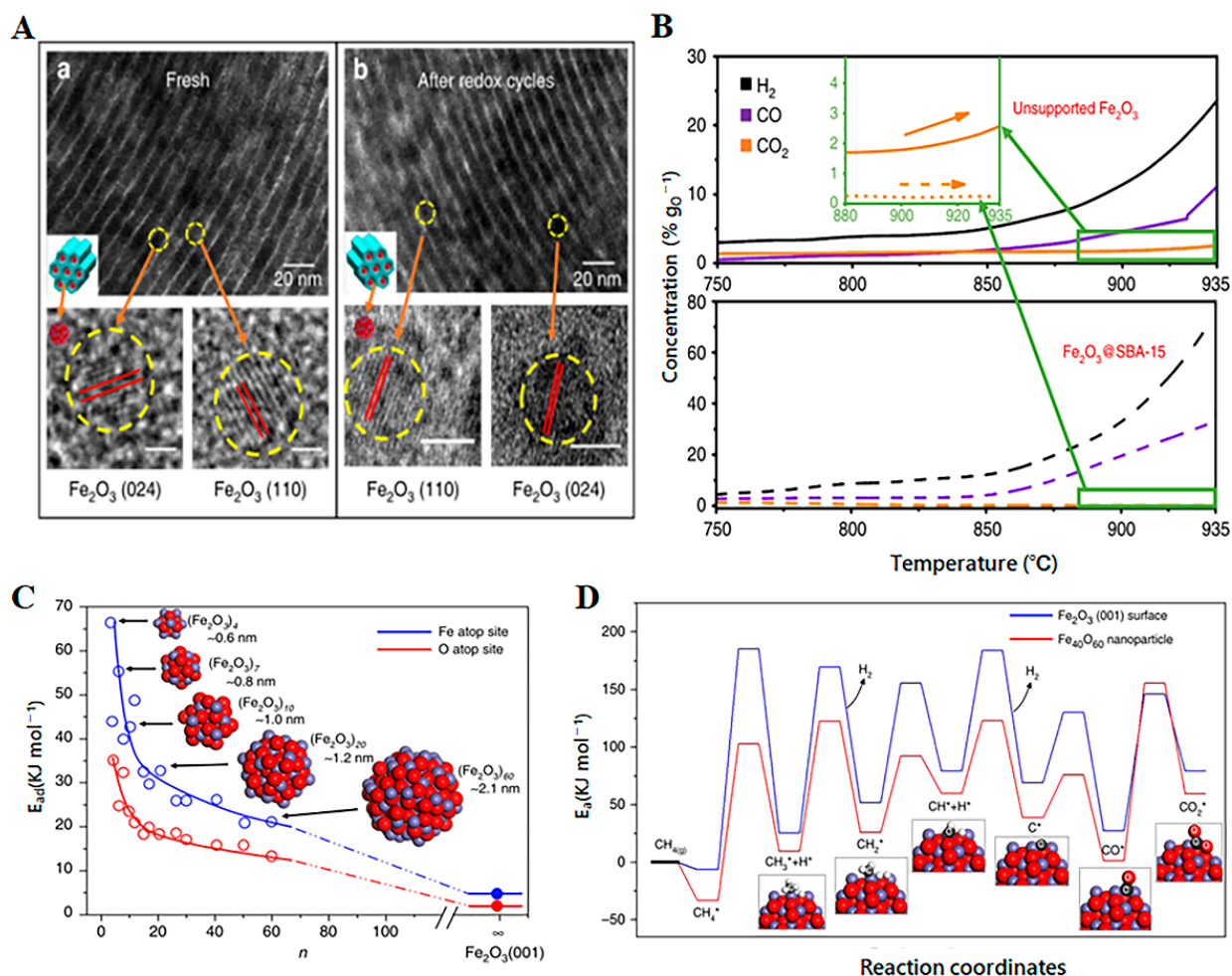


Figure 7. (A) Morphological characteristics of $\text{Fe}_2\text{O}_3/\text{SBA-15}$ OC. (Aa) HR-TEM images of fresh $\text{Fe}_2\text{O}_3/\text{SBA-15}$ and typical Fe_2O_3 nanoparticles (scale bar of 1 nm); (Ab) HR-TEM images of $\text{Fe}_2\text{O}_3/\text{SBA-15}$ after 75 cycles and typical Fe_2O_3 nanoparticles (scale bar of 5 nm); (B) Temperature programmed reduction profiles of $\text{Fe}_2\text{O}_3/\text{SBA-15}$ and unsupported Fe_2O_3 ; (C) Calculated results of CH_4 adsorption on Fe atop site and O atop site of $(\text{Fe}_2\text{O}_3)_n$ nanoparticles; (D) Energy evolution of CH_4 conversion on $\text{Fe}_{40}\text{O}_{60}$ nanoparticle and Fe_2O_3 (001) surface. Adapted with permission from Ref. [102], copyright 2019, Nature.

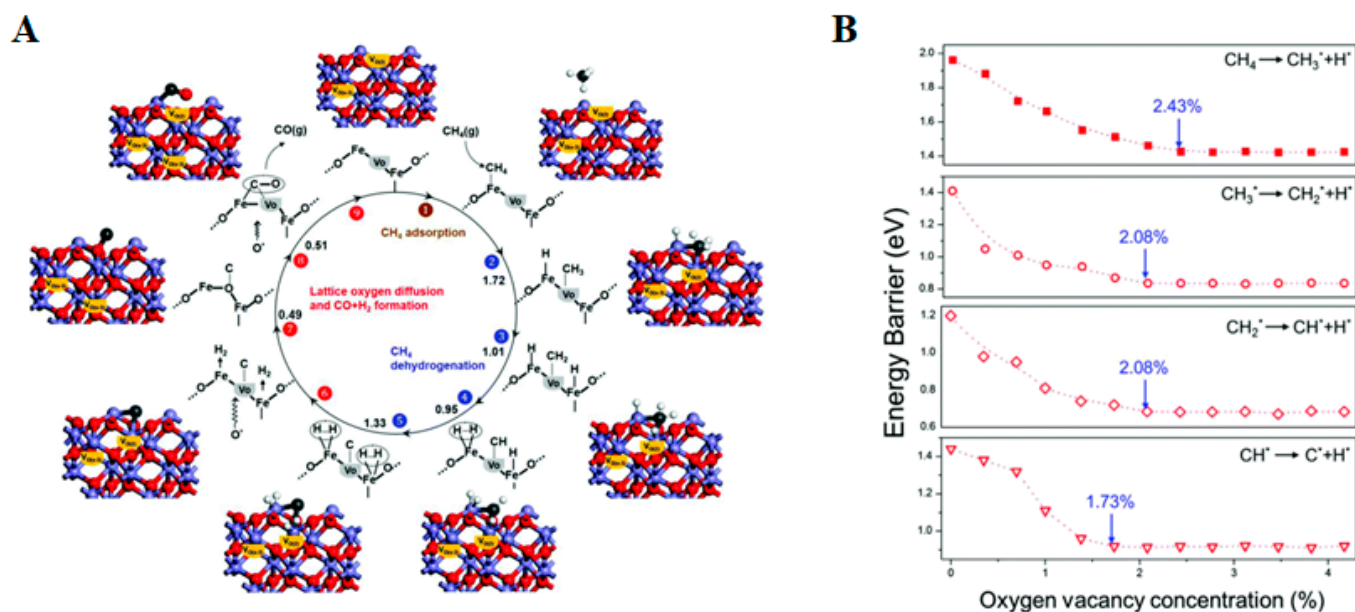


Figure 8. (A) Mechanism for CH₄ partial oxidation over α-Fe₂O₃ (001) surface with oxygen vacancies. Inset of the reaction loop is the activation barrier (eV) for each elementary step. V_O represents the surface oxygen vacancy. (B) The correlation between activation energy of C–H bond and the oxygen vacancy concentration of α-Fe₂O₃ (001) surface. Reprinted with permission from Ref. [103], copyright 2016, Royal Society of Chemistry.

2.2.2. Supported or Doped Cerium Oxides

Cerium-based OCs have been widely studied for CL-SMR in terms of their high oxygen storage capacity and reliable resistance to carbon deposition [105–107]. However, pure CeO₂ exhibits low reactivity towards methane conversion and is prone to sintering in several cyclic reactions. Therefore, great effort has been applied to improve the reactivity and stability of these OCs by doping foreign cations and constructing composite oxides. Wang et al. [108] discovered that Ni-modified (5 wt%) CeO₂-TiO₂ enabled 100% conversion of CH₄, ca. 16 times greater than that of pure CeO₂-TiO₂ (5.9%), with 85% syngas selectivity at 900 °C. A mechanism study showed that Ni species accounted for the main active centers for CH₄ activation and promoted the reduction of Ce⁴⁺ (CeO₂-TiO₂) to Ce³⁺ (Ce₂Ti₂O₇) (Figure 9B). The synergy between active Ni species and CeO₂-TiO₂/Ce₂Ti₂O₇ redox oxides significantly increased the water-splitting performance with hydrogen yield up to 47.0 mL/g (Figure 9). Zhu et al. [87,109,110] prepared a series of Fe-doped cerium oxides (Ce_{1-x}Fe_xO_{2-δ}) for a CL-SMR process and discovered that introduction of iron notably improved the reducibility of OCs, and formation of Ce-Fe-O solid solution (CeFeO₃) could enhance the lattice oxygen migration, leading to increased methane conversion. As for the optimized sample (x = 0.5), conversion of methane approached 60% with syngas selectivity exceeding 80% and hydrogen productivity of 83.3 mL/g. Li et al. [111] loaded Ce-Fe-Zr-O oxides (40 wt%) onto inert MgAl-LDO (magnesium–aluminum layered double oxides) and demonstrated that the redox reactivity and stability were greatly improved due to the highly dispersed active components, accelerated reaction between Ce-Fe oxides, and restrained sintering of OCs. Experimental results showed that methane conversion decreased from 58% to 46% in the first 6 cycles and stayed steady in the following 14 cycles with CO selectivity maintained over 80% and hydrogen yield of 1.0 mmol/g. This group further studied the synergy between CeO₂ and 3DOM-LaFeO₃ during CL-SMR reactions and found that 10 wt% CeO₂/LaFeO₃ displayed the highest yields of syngas (9.94 mmol/g) and hydrogen (3.38 mmol/g), since the interaction of CeO₂-LaFeO₃ generated a great number of oxygen vacancies that promoted lattice oxygen diffusion and restricted the carbon deposition [112]. Zhu et al. [113] also found an obvious synergistic effect between CeO₂ and BaFe₃Al₉O₁₉ (BF3). The difference was that a new phase of CeFe_xAl_{1-x}O₃ was generated

during chemical looping reactions, which resulted in promoted diffusion of lattice oxygen and increased methane conversion to ca. 90%. Wei et al. [114] investigated the influence of OCs morphology on the redox reactivity and found that $\text{CeO}_2/\text{ZrO}_2$ oxides calcined at low temperatures (450°C) held stable 3DOM structure with uniform particle size, which effectively enhanced the migration of oxygen species and improved the reactivity for CH_4 partial oxidation and water-splitting. After 10 cycles at 800°C , a hydrogen yield exceeding 38.89 mL/g was maintained.

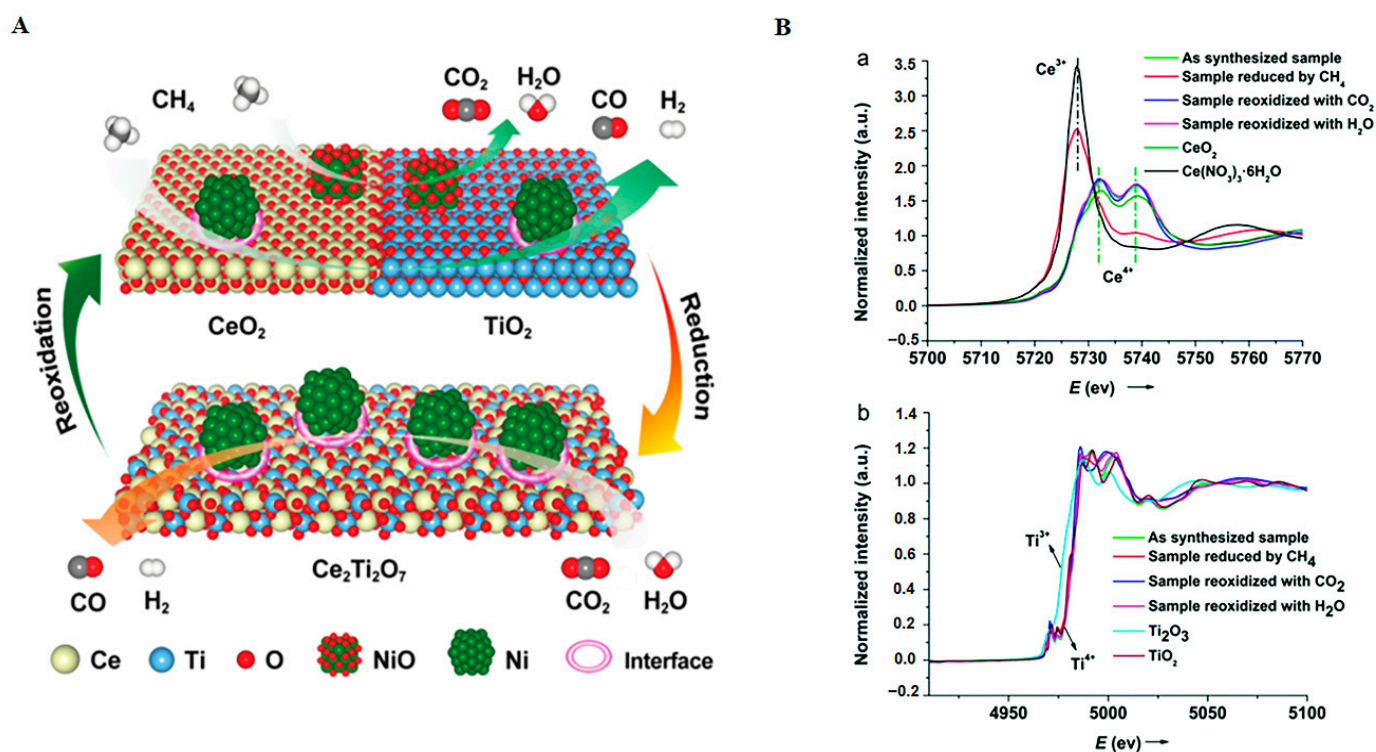


Figure 9. (A) Proposed mechanism for MDR-STCDS and MDR-STWS processes over $5\text{Ni}/\text{CeO}_2\text{-TiO}_2$. (B) Normalized XANES profiles for $5\text{Ni}/\text{CeO}_2\text{-TiO}_2$ after various steps during isothermal MDR-STCDS and MDR-STWS reactions at (Ba) the Ce L_{III}-edge and (Bb) the Ti K-edge, together with the spectra of reference compounds. Adapted with permission from Ref. [108], copyright 2019, Royal Society of Chemistry.

2.2.3. Perovskites

Perovskite oxides are promising OCs for CL-SMR due to their excellent redox activity [115,116]. Lee et al. [117] discovered that doping Fe in B-site of LaCoO_3 improved lattice oxygen mobility, reactivity for partial oxidation of CH_4 , and adsorption and dissociation of H_2O , which enabled high CO selectivity of 92% and H_2 purity of 99.3% during the methane reduction and water-splitting steps, respectively. Gong and colleagues [118,119] discovered that substitution of Ce^{3+} for La^{3+} of LaFeO_3 was capable of tuning lattice oxygen activity via modulating the distortion degree of FeO_6 octahedra. When Ce/La ratio reached 1 ($\text{La}_{0.5}\text{Ce}_{0.5}\text{FeO}_3$), the oxygen carrier had the lowest oxygen vacancy formation energy and the highest oxygen mobility, which rendered the best methane-to-syngas and water-splitting performance. Furthermore, the yield of hydrogen (0.6 mmol/g) remained stable during 100 redox cycles (Figure 10), highlighting the outstanding stability of this OC. Recently, Wang et al. [120] proposed that the reactivity of lattice oxygen in $\text{LaFe}_{0.8}\text{M}_{0.2}\text{O}_3$ ($\text{M} = \text{Al, Fe, Ga, Sc}$) perovskite was not only influenced by B-site cations, but also greatly affected by the La cations at A-site. It was found that a decline of FeO_6 tilting degree by doping smaller cations at B-site of $\text{LaFe}_{0.8}\text{M}_{0.2}\text{O}_3$ was beneficial to weaken the La–O interaction, thereby improving the mobility of lattice oxygen and methane conversion performance. The researchers [121] further showed that doping Sr and Al at A and B sites

of LaFeO_3 , respectively, significantly promoted the deep reduction of $\text{Fe}^{4+/3+}$ cations to Fe^0 by forming Fe^0 @oxides intermediate with core-shell structure. Moreover, the core-shell structure formed in methane atmosphere was able to prevent direct contact between CH_4 and Fe^0 , which promoted syngas production and suppressed coke deposition over Fe^0 (Figure 11). Li's group [122] discovered that doping of Fe in LaMnO_3 greatly enhanced the redox performance due to lattice distortion and increased concentration of oxygen vacancy. Among the screened oxides, $\text{La}_{0.85}\text{MnFe}_{0.15}\text{O}_3$ displayed the highest CO selectivity of ~99% and hydrogen yield of 1.76 mmol/g. Similar results could be observed for Ni or Co modified LaMnO_3 [123]. Li et al. [124] designed an OCs of Fe_2O_3 @ $\text{La}_x\text{Sr}_{1-x}\text{FeO}_3$ (LSF) with core-shell structure and found that LSF not only improved the diffusion of lattice oxygen in Fe_2O_3 but also reduced the direct contact between CH_4 and Fe_2O_3 , which resulted in highly selective CH_4 -to-syngas conversion with selectivity as high as 89%. Zhao et al. [125] found that $\text{LaFe}_{1-x}\text{Co}_x\text{O}_3$ ($x = 0, 0.2, 0.4, 0.6, 0.8$) with a 3DOM structure promoted the mass transfer and provided more active centers for CH_4 conversion. Moreover, Co doping increased oxygen mobility, which reduced carbon deposition and improved the purity of hydrogen. This group [126] further investigated the synergistic effect of different metals in double perovskite $\text{La}_{1.6}\text{Sr}_{0.4}\text{FeCoO}_6$, and found that CH_4 was prone to be activated on cations with high valence state (Fe^{5+} , Fe^{4+} , and Co^{3+}), while the correspondingly formed Fe^{2+} , Co^0 and the oxygen vacancies provided rich active centers for splitting of H_2O .

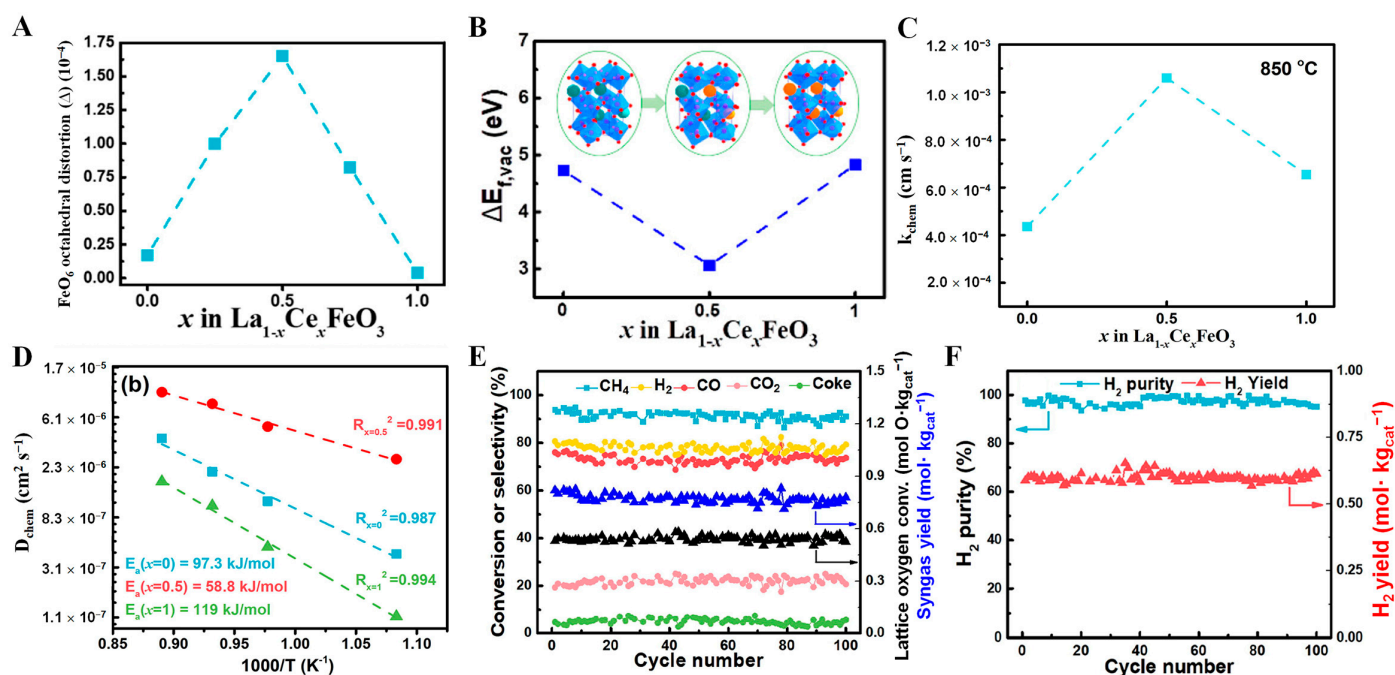


Figure 10. (A) Distortion degree (Δ) of FeO_6 octahedra for $\text{La}_{1-x}\text{Ce}_x\text{FeO}_3$ ($x = 0, 0.25, 0.5, 0.75, 1$) and (B) calculated oxygen vacancy formation energy ($\Delta E_{f,vac}$) of $\text{La}_{1-x}\text{Ce}_x\text{FeO}_3$ ($x = 0, 0.5, 1$). Adapted with permission from Ref. [119], copyright 2020, American Chemical Society. (C) The oxygen surface exchange coefficient (k_{chem}) and (D) bulk diffusion coefficient (D_{chem}) of $\text{La}_{1-x}\text{Ce}_x\text{FeO}_3$ ($x = 0, 0.5, 1$). Redox performance of (E) CH_4 partial oxidation step and (F) the H_2O -splitting step over $\text{La}_5\text{Ce}_5\text{FeO}_3$ in CL-SRM at 925°C . Adapted with permission from Ref. [118], copyright 2020, Elsevier.

2.3. Chemical Looping Water Gas Shift Process

As a low-cost but important platform chemical, CO can be used as a reductant-like methane to drive the CLWS process and produce high-value hydrogen (Figure 2C) [127,128]. Herein, OCs were reduced by CO ($\text{CO} + \text{O}_L \rightarrow \text{CO}_2$) and subsequently regenerated by water ($\text{H}_2\text{O} \rightarrow \text{O}_L + \text{H}_2$). This process is generally known as a chemical looping water gas shift (CL-WGS) reaction due to the same overall reaction as that of water-gas shift reaction ($\text{CO} + \text{H}_2\text{O} \rightarrow \text{CO}_2 + \text{H}_2$). Compared with CL-SMR, the application of CO as a

reducing atmosphere can avoid the carbon deposition caused by CH_4 decomposition while the coke formation from the Boudouard reaction ($2\text{CO} \rightarrow \text{C} + \text{CO}_2$) is thermodynamically constrained at a temperature above 800°C , which inhibited the contamination of hydrogen by carbon monoxide ($\text{C} + \text{H}_2\text{O} \rightarrow \text{CO} + \text{H}_2$). Furthermore, the separation of CO/CO_2 from $\text{H}_2\text{O}/\text{H}_2$ in chemical looping reactions also prevents the reverse water–gas shift reaction ($\text{CO}_2 + \text{H}_2 \rightarrow \text{CO} + \text{H}_2\text{O}$), leading to increased efficiency for water-splitting. Thermodynamic analysis demonstrated that the steam conversion in CL-WGS reaction reached 95% at 800°C , which is significantly higher than the traditional WGS process [32]. Based on the ASPEN Plus simulation calculation, Zeng et al. [129] found that the thermal efficiency of H_2 production by a chemical looping process, including reduction of OCs by coal and CO (reducer reactor), partial oxidation of OC by steam oxidation to produce hydrogen (oxidizer reactor), and completely recovery of OC by air (combustor), could reach 78% for HHV (higher heating value) efficiency, which was much higher than that of hydrogen production from the traditional coal gasification method.

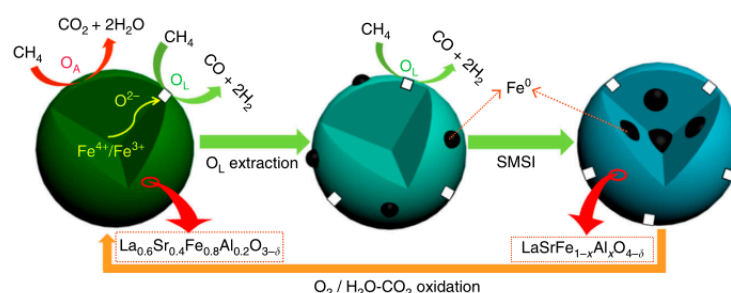


Figure 11. Proposed structural evolution of $\text{La}_{0.6}\text{Sr}_{0.4}\text{Fe}_{0.8}\text{Al}_{0.2}\text{O}_{3-\delta}$ OC during chemical looping partial oxidation of methane. White boxes represent the coordination unsaturated Fe cations. Reprinted with permission from Ref. [121], copyright 2018, Nature.

As for CL-WGS, the suitable OCs should be reactive for CO oxidation and subsequent water-splitting while possessing high redox stability. Fe_2O_3 has been widely studied due to the features of abundant reserves, favorable thermodynamic properties, and high oxygen-carrying capacity [130]. However, Fe_2O_3 is prone to agglomerate during several redox cycles, rendering rapid decline of hydrogen productivity. Recent works showed that loading of Fe_2O_3 on some inert supports, including ZrO_2 [131], Al_2O_3 [132,133], and MgAl_2O_4 [134], were effective in improving the specific surface area of OCs and slowing down the sintering process. Ma et al. [135] investigated in detail the effects of redox inert supports on the reactivity, redox stability, and hydrogen yield of Fe_2O_3 /support OCs (support = Al_2O_3 , SiO_2 , MgAl_2O_4 , and ZrO_2). It was found that the hydrogen yield of $\text{Fe}_2\text{O}_3/\text{Al}_2\text{O}_3$ and $\text{Fe}_2\text{O}_3/\text{SiO}_2$ gradually decreased with cyclic numbers, since solid phase reaction between Al_2O_3 or SiO_2 and Fe_2O_3 induced formation of less reactive FeAl_2O_4 and FeSiO_3 , respectively. In contrast, when MgAl_2O_4 was utilized, the solid phase interaction between Fe_2O_3 and support was greatly inhibited, leading to improved redox stability and hydrogen yield of 54–93 mL/g. Yuzbasi et al. [136] reported that Na-modified $\text{Fe}_2\text{O}_3/\text{Al}_2\text{O}_3$ could generate a layered Na- β - Al_2O_3 phase to restrict the formation of FeAl_2O_4 , which increased the redox stability and rendered a stable hydrogen production of 13.3 mmol/g during 15 cycles. Compared with these inert supports, introduction of MIEC support showed a much-pronounced effect in improving the efficiency of hydrogen production by enhancing migration of lattice oxygen and electron conductivity [137]. Ma et al. [138] studied the performance of $\text{A}_x\text{Ce}_{1-x}\text{O}_{2-\delta}$ ($\text{A} = \text{Y}, \text{Sm}$ and La) supported Fe_2O_3 for CL-WGS and found that Sm doping ($\text{Fe}_2\text{O}_3/\text{Ce}_{0.7}\text{Sm}_{0.2}\text{O}_{1.9}$) exhibited the best oxygen anion conductivity and oxygen mobility, which facilitated the conversion of CO and subsequent hydrogen production. In contrast, the reactivity of OCs was negatively correlated with La content due to the decreased oxygen capacity arising from formation of LaFeO_3 during the reaction. They further discovered that Zr and Sm co-doped $\text{Fe}_2\text{O}_3/\text{Ce}_{0.8}\text{A}_{0.2}\text{O}_{2-\delta}$ ($\text{A} = \text{Zr}$

and Sm) displayed better CO conversion and sintering resistance, rendering a hydrogen yield of 54.25 mL/g for water-splitting [139].

Iron spinels that bear features of low cost, high thermal stability, and good oxygen mobility are promising OCs for chemical looping reactions [140,141]. Huang et al. [142] found that $\text{NiFe}_2\text{O}_4/\text{Al}_2\text{O}_3$ displayed stable H_2 yield during 20 cycles of CL-WGS, since Al_2O_3 support significantly inhibited the sintering of NiFe_2O_4 . Kim et al. [143] also found that Al-modified NiFe_2O_4 exhibited good performance for water-splitting due to the formation of a spinel solid solution with Al, which could prevent densification of NiFe_2O_4 . For the optimized sample with Al content of 3.3 wt%, maximum hydrogen productivity of 8.2 mmol/g was reached. Cui et al. [144] reported that the active Fe could completely exsolve from mixed Zn-Fe-Al spinel in CO atmosphere and resolved into spinel structure by water oxidation, enabling a hydrogen yield of 2.23 mmol/g with nearly 100% conversion of CO. More importantly, an embedded interface structure was formed for the reduced OC, which effectively limited Fe agglomeration and improved the reactivity and stability of the OCs. Based on DFT+U calculations, Xiao et al. [145] demonstrated that the reactivity of binary spinel AFe_2O_4 ($\text{A} = \text{Co}, \text{Cu}, \text{Ni}, \text{Mn}$) was mainly related to the concentration of oxygen vacancies (reduction level of OCs). Among these OCs, Co-modified OC was the best candidate for CL-WGS due to its suitable oxygen vacancy formation energy (2.70 eV) and lattice oxygen diffusion energy (1.03 eV), which gave an impressive yield of 11.9 mmol/g with average hydrogen production rate of 0.051 mmol/g·s at 650 °C (Figure 12). They further discovered that the oxygen vacancy formation energy and oxygen diffusion energy further decreased to 1.69 eV and 0.83 eV, respectively, after further doping of Cu in CoFe_2O_4 ($\text{Cu}_{0.25}\text{Co}_{0.25}\text{Fe}_{2.5}\text{O}_4$), which promoted the reduction kinetics in CO environment and increased the redox efficiency for hydrogen production. As a result, a hydrogen production as high as ~11.9 mmol/g was achieved at a temperature of 550 °C [146].

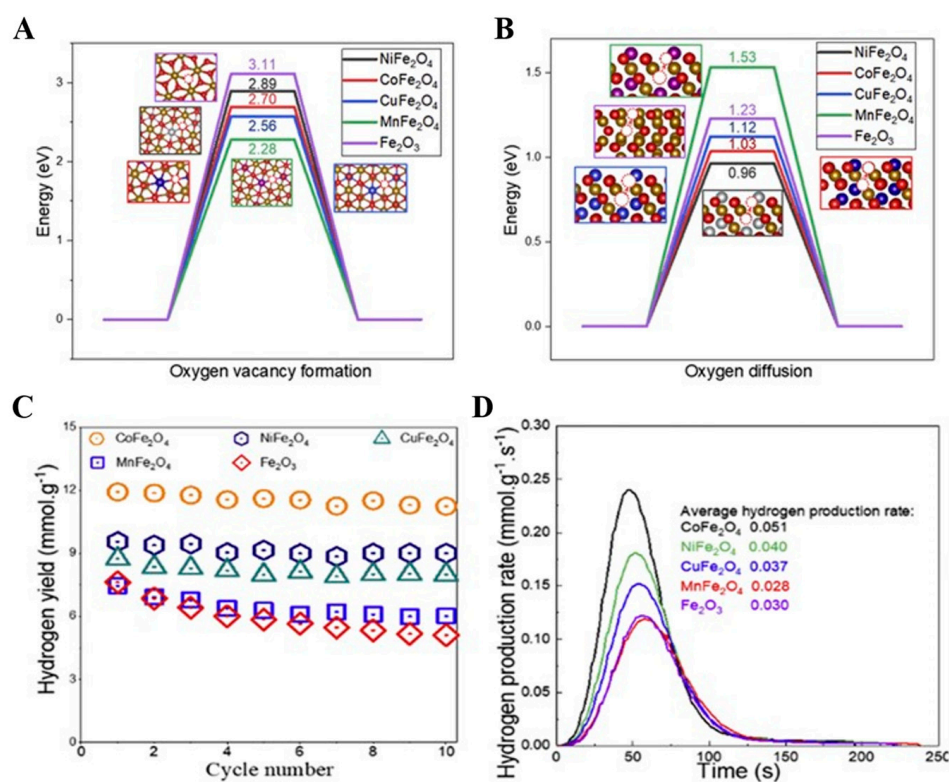


Figure 12. (A) The oxygen vacancy formation energy and (B) diffusion energy of lattice oxygen in CoFe_2O_4 (100), CuFe_2O_4 (100), NiFe_2O_4 (100), MnFe_2O_4 (100), and Fe_2O_3 (001) surfaces. O, Cu, Co, Ni, Mn, and Fe atoms are shown with red, light blue, dark blue, light grey, purple, golden spheres, respectively. (C) Profiles of the activation energy and (D) the hydrogen yield for different OCs. Reprinted with permission from Ref. [145], copyright 2019, Elsevier.

Brownmillerite ($\text{Ca}_2\text{Fe}_2\text{O}_5$) is well known for high oxygen capacity and excellent thermal stability due to the anion-deficient structure with alternating FeO_6 octahedral and FeO_4 tetrahedral layers [147,148]. Based on the thermodynamic study, Ismail et al. [149] found that the valence state of Fe in $\text{Ca}_2\text{Fe}_2\text{O}_5$ only displayed Fe^0 and Fe^{3+} without intermediate Fe^{2+} during the redox reactions, and Fe^0 could be directly oxidized to Fe^{3+} by steam, which was suitable for a water-splitting reaction. Chan et al. [150] discovered that the steam conversion over $\text{Ca}_2\text{Fe}_2\text{O}_5$ could reach 75%, which was much higher than that of $\text{Fe}_2\text{O}_3/\text{ZrO}_2$ (62%). Through thermogravimetric experiments, Sun et al. [151] found that $\text{Ca}_2\text{Fe}_2\text{O}_5$ displayed much faster reaction kinetics than CaFe_2O_4 during the CO reduction step, rendering a higher reduction degree of 94.0% (85.5% for CaFe_2O_4). Guo et al. [152] employed DFT calculations to explore the mechanisms of CO adsorption, oxidation, and oxygen anion diffusion on the surface of $\text{Ca}_2\text{Fe}_2\text{O}_5$ (010). It was discovered that CO preferred to adsorb on FeO_2 - and FeO -terminated surfaces rather than CaO - and OCa -terminated surfaces, while the chemisorption energy of FeO_2 -terminated surfaces was considerably greater than that of FeO -terminated surfaces. In addition, it is also proposed that the oxygen anion diffusion at high temperature was the rate-limiting step because the energy barrier of oxygen diffusion was significantly greater than that of the CO oxidation process (Figure 13).

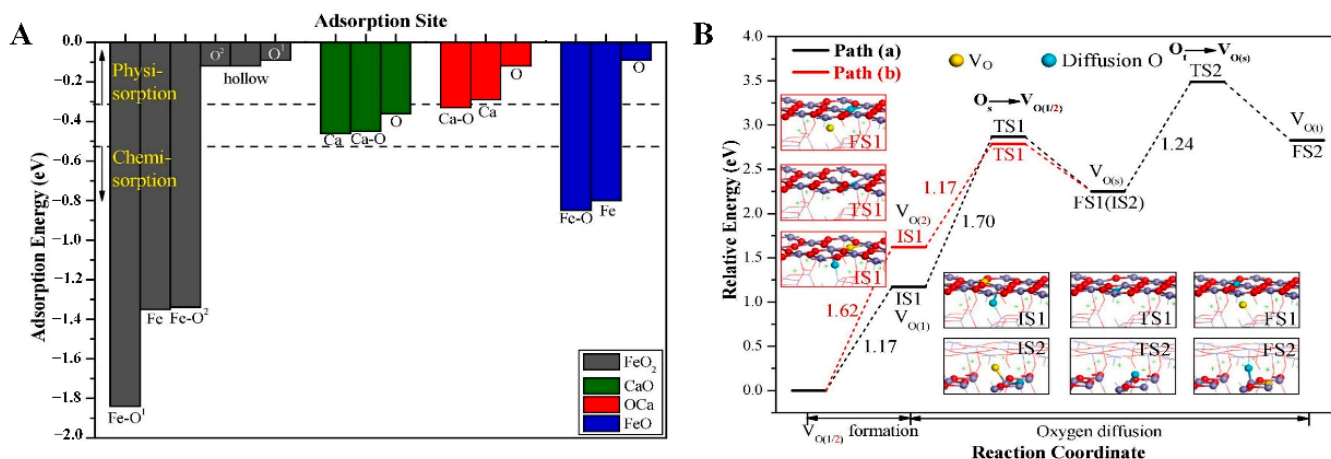


Figure 13. (A) Comparison of CO adsorption energies on FeO_2 -, CaO -, FeO -, and OCa -terminated $\text{Ca}_2\text{Fe}_2\text{O}_5$ (0 1 0) surfaces. (B) The pathways and energy profiles for oxygen diffusion in $\text{Ca}_2\text{Fe}_2\text{O}_5$. Adapted with permission from Ref. [152], copyright 2020, Elsevier.

2.4. Syngas Chemical Looping Process

Syngas that can be produced via various routes, e.g., gasification of biomass, coal, and methane reforming, and represents another promising reducing agent for CLWS [153–155]. In the syngas-promoted CLWS process (Figure 2D), referred to as syngas chemical looping (SCL), syngas was utilized as the reducing gas instead of CO, while subsequent water-splitting and air oxidation (in some cases) were needed to recover the OCs. The focused OCs explored for this process are mainly iron-based oxides.

Fan et al. [33] investigated the feasibility of Ni, Cu, Cd, Co, Mn, Sn, and Fe oxides for SCL through thermodynamic analysis, and found that Fe_2O_3 exhibited the best syngas and steam conversion. The Fe_2O_3 OC displayed stable redox performance during 10 cycles at 600 °C and possessed a low attrition rate of 0.57% during the redox process in entrained flow reactor [156]. When a moving bed reactor was utilized, they found that a syngas conversion of 99.95% with Fe_2O_3 conversion close to 50% was achieved at 900 °C [157]. Dou et al. [158] showed that the syngas generated by in situ decomposition of glycerol was effective for reduction of OCs. Among the investigated OCs with different Ce/Ni/Fe molar ratios (0/0/1; 10/1/100, 10/3/100, 5/1/100, 5/2/100), OC with Ce/Ni/Fe ratio of 10/3/100 displayed the highest H_2 yield of 11.79 mmol/g at 750 °C. Mechanism stud-

ies suggested that Ni dopants promoted glycerol conversion while Ce could effectively improve the dispersion of Ni, the sintering resistance, and lattice oxygen mobility, thus substantially boosting the hydrogen yield. Sun et al. [159] investigated the performance of CeO₂-modified Fe₂O₃/Al₂O₃ in SCL process and found that CeO₂ could coordinate with Fe to generate more oxygen vacancies, which promoted the participation of lattice oxygen in the reaction and inhibited the formation of carbon or Fe₃C. The best OC with Fe₂O₃/CeO₂/Al₂O₃ ratio of 65/5/30 OCs achieved the best hydrogen productivity, while the overall performance only decreased by 21% during eighth cycles.

Based on thermodynamic analysis and packed bed experiments, Aston et al. [160] found that the conversion of syngas reached more than 99% during the reduction process of NiFe₂O₄ and CoFe₂O₄. Furthermore, the reduced NiFe₂O₄ (NiFe alloy) and CoFe₂O₄ (CoFe alloy) was highly reactive towards water-splitting with regeneration of spinel structure, which rendered the NiFe₂O₄ and CoFe₂O₄, bearing better redox performance than Fe₂O₃ (not recovered by water) for SCL process. He et al. [161] investigated the effect of preparation methods, including solid-state method, coprecipitation method, hydrothermal method, and sol-gel method, on the SCL performance of NiFe₂O₄ nanoparticles. It was found that the OC prepared by sol-gel method displayed the best hydrogen yield and highest lattice oxygen recovery degree due to its smaller particle size and porous structure. Furthermore, further mechanical mixing of NiFe₂O₄ and inert SiO₂ could effectively restrain the sintering of NiFe₂O₄, rendering outstanding redox stability at 850 °C for 20 cycles. Scheffe et al. [162] prepared m-ZrO₂ (surface area of 50 m²/g) supported Co_{0.85}Fe_{2.15}O₄ via atomic layer deposition (ALD) method and found that such OC produced much more hydrogen than that over γ-Fe₂O₃/m-ZrO₂ in 7 cycles. Mechanism studies suggested that the high yield of hydrogen was ascribed to the deep reduction of Fe/Co cations into CoFe alloy, exceeding the reduction degree of γ-Fe₂O₃ (γ-Fe₂O₃ → FeO), which improved the oxygen capacity and water-splitting reactivity, and thus, the hydrogen productivity (Figure 14).

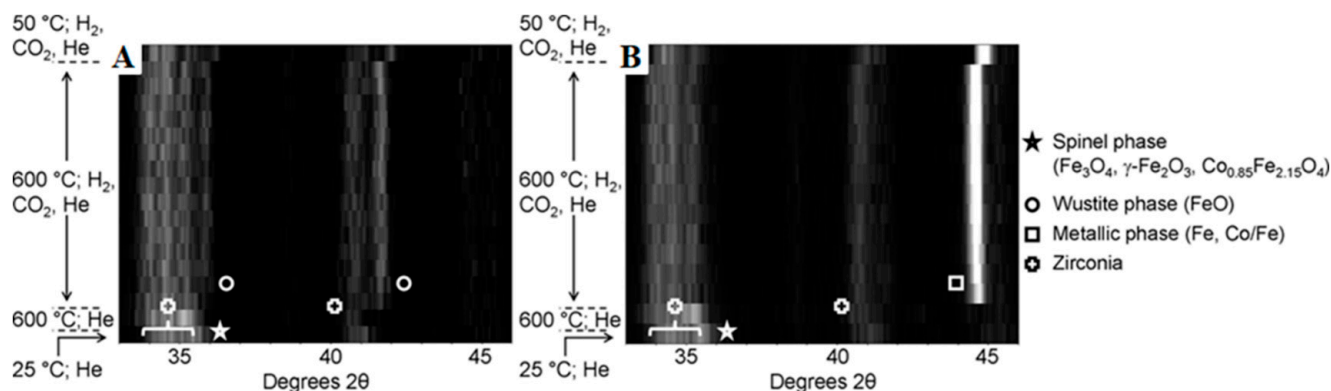


Figure 14. In situ XRD profiles of OCs reduced by 1 vol % H₂/1 vol % CO₂ in He at 600 °C. (A) Fe₂O₃ supported on m-ZrO₂ (ALD/20.2% mass loading) and (B) Co_{0.85}Fe_{2.15}O₄ supported on m-ZrO₂ (ALD/19% mass loading). Chemical reduction for 148 min. Adapted with permission from Ref. [162], copyright 2011, American Chemical Society.

2.5. Photo-Thermochemical Cycle

In 2016, Zhang and coworkers [34] first proposed a strategy of photo-thermochemical cycle (PTC) for CLWS, which explored the photochemical reduction method instead of thermal reduction. This method not only considerably decreased the threshold for abstracting lattice oxygen from OC, slowed down the sintering process, and improved the cyclic stability, but also enabled transformation of solar energy into chemical energy. In a typical photo-thermochemical cycle for water-splitting, the OCs were firstly reduced using ultraviolet and visible light at room temperature with releasing oxygen to generate photo-induced oxygen vacancies, and subsequently recovered by water-splitting via infrared heating to temperature of 500~600 °C with production of hydrogen.

Up to now, the most studied OCs for PTC have been restricted to pure or modified TiO₂ photocatalysts. In 2016, TiO₂ was used as OC for PTC water-splitting to produce hydrogen, and the average hydrogen production of each cycle reached 0.421 mL/g in five consecutive cyclic reactions [34]. However, TiO₂ has a narrow photoresponse range, high recombination rate of photo-induced electron-hole pairs (EHPs), and low reactivity with water. To address these problems, Xu et al. [163] prepared 0.5 wt% Fe/TiO₂ as an OC, which displayed much higher hydrogen yield of 33.36×10^{-3} mmol/g than TiO₂ (18.79×10^{-3} mmol/g). This is ascribed to the lower recombination rate of EHPs (Figure 15A) and wider photoresponse range (Figure 15B) after doping Fe (III) cations. Further DFT calculations showed that introduction of Fe cations could reduce the oxygen vacancy formation energy, improve H₂O adsorption, and enhance the desorption of generated H₂, which substantially boosted the efficiency of H₂ production (Figure 15C). Docao et al. [164] reported that doping Cu in TiO₂ could raise the distortion degree of TiO₆ octahedron, which improved the oxygen mobility and promoted water-splitting at 140 °C, with solar energy conversion efficiency reaching 0.63%.

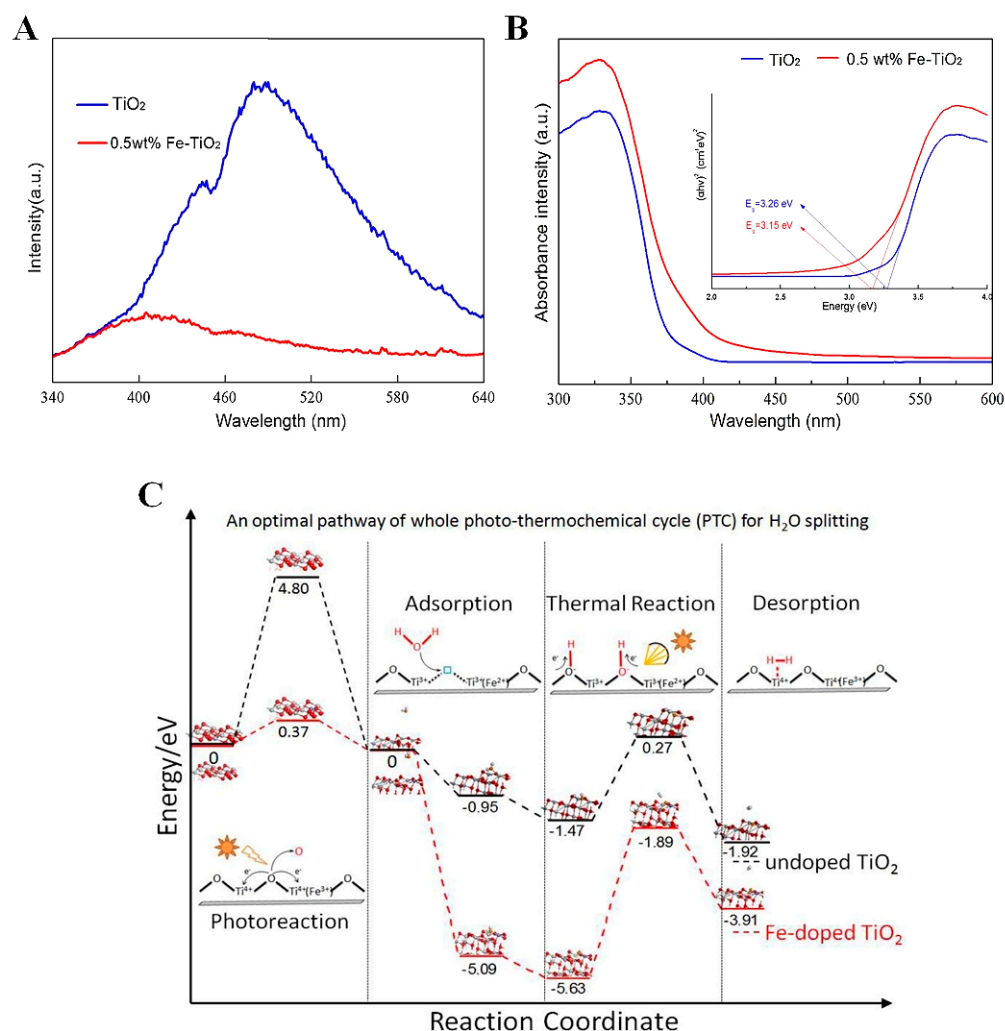


Figure 15. (A) PL profiles for TiO₂ and 0.5 wt% Fe-doped TiO₂. (B) UV-vis DRS profiles for TiO₂ and 0.5 wt% Fe-doped TiO₂. Inset shows the optical energy band gap. (C) The reaction pathway of H₂O splitting during PTC process. Reprinted with permission from Ref. [163], copyright 2017, Elsevier.

Wu et al. [165] modified TiO₂ OCs by doping various transition metals, including Fe, Cu, Co, Ni, and Zn, and discovered that these modified OCs showed higher oxygen vacancy concentration and wider photoresponse range than bare TiO₂. Among these OCs, Cu-doped TiO₂ displayed the highest light absorption range and the lowest recombination

rate of EHPs, which notably improved the driving force for the release of lattice oxygen and hydrogen production. Further decoration of Cu/TiO₂ by Ni could generate more oxygen vacancies and enhance the activity for water-splitting, which exhibited the maximum hydrogen output of 27.01×10^{-3} mmol/g, nearly 36.0 times higher than that of undoped TiO₂ (0.75×10^{-3} mmol/g) (Figure 16). The follow-up research showed that 0.1~2.0 wt% of Ni-promoted Cu/TiO₂ could anchor nickel oxides (Ni₂O₃ and NiO) around the surface oxygen vacancy while the low valent state NiO showed more pronounced promotion effect towards water-splitting, which rendered the 0.5 wt% Ni-promoted Cu/TiO₂ with highest NiO content, displaying a hydrogen yield (30.6×10^{-3} mmol/g) 40 times higher than that of undoped TiO₂ [166].

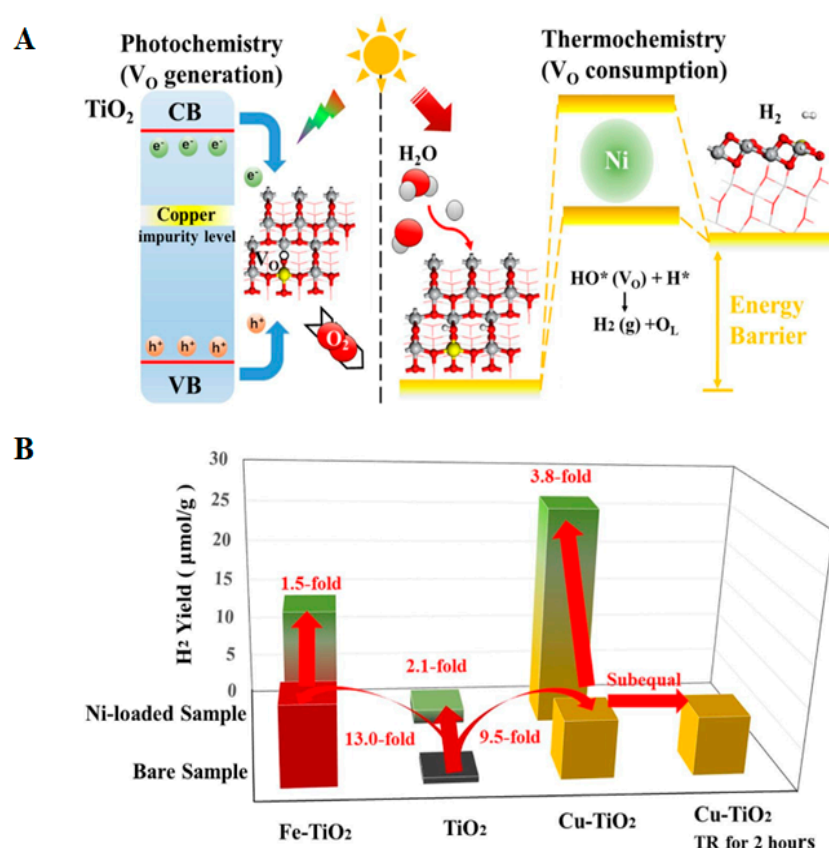


Figure 16. (A) Schematic of the reaction mechanism for photochemical and thermochemical processes. (B) The average H₂ yield of different catalysts. TR represented “thermochemical reaction”. Reprinted with permission from Ref. [165], copyright 2021, Elsevier.

3. Summary and Outlook

Chemical looping water-splitting is promising for sustainable hydrogen production due to the virtue of decoupling a one-step reaction into two or three spatially separated reactions, which greatly simplifies the gas separation process and avoids the harsh conditions for direct water decomposition. To date, different processes have been developed, including a two-step thermochemical water-splitting (TCWS) cycle, methane chemical looping process, chemical looping water gas shift (CL-WGS) cycle, syngas chemical looping (SCL) process, and photo-thermochemical cycle (PTC), with attempts to reduce the energy penalty and CO₂ emissions by altering the method to abstract the lattice oxygen from the OCs, wherein the key lies in the manufacture of suitable OCs. Over the past decades, thousands of metal oxides, such as iron-based metal oxides, zinc-based metal oxides, cerium-based metal oxides, titanium-based metal oxides, and composite metal oxides (perovskite, spinel and garnets, etc.), have been prepared for chemical looping reactions. The performance summary of OCs in different processes of CLWS is summarized in Table 1.

Table 1. Performance summary of OCs in different processes in CLWS.

Hydrogen Production Processes	OCs Type	Cyclic Reaction	OCs	Reaction Temperature (°C; Red/Ox)	H ₂ Yield (mmol/g)	Ref.
Two-step TCWS cycle	Ferrites (M _x Fe _{3-x} O ₄)	$M_xFe_{3-x}O_4 \rightarrow xMO + (3-x)FeO + 0.5O_2$; $xMO + (3-x)FeO + H_2O \rightarrow M_xFe_{3-x}O_4 + H_2$	NiFe ₂ O ₄	1200/800	0.0197 *	[48]
			Co _{0.67} Fe _{2.33} O ₄ /ZrO ₂	1300~1400/1000	0.357~0.535	[49]
			NiFe ₂ O ₄	1400/1000	~0.446	[50]
	ZnO ₂	ZnO → Zn + O ₂ ; Zn + H ₂ O → ZnO + H ₂	NiFe ₂ O ₄ /m-ZrO ₂	1400/1000	0.535~0.714	[50]
			ZnO	-	-	
	Cerium oxide (Ce _x M _{1-x} O ₂)	$Ce_xM_{1-x}O_2 \rightarrow Ce_xM_{1-x}O_{2-\delta} + 0.5\delta O_2$; $Ce_xM_{1-x}O_{2-\delta} + H_2O \rightarrow \delta H_2O \rightarrow Ce_xM_{1-x}O_2 + \delta H_2$	CeO ₂	1350/800	0.032	[31]
			CeO ₂	1500/500	0.142 *	[66]
			Ce _{0.9} Hf _{0.1} O ₂	1500/500	0.2 *	[66]
			Ce _{0.9} Sc _{0.1} O _{1.95}	1500/500	0.181 *	[66]
			Ce _{0.75} Zr _{0.25} O ₂	1400/1050	0.297	[69]
			1% Gd/CeO ₂ -ZrO ₂	1400/1050	0.338 *	[71]
			CeO ₂ -0.15SnO ₂	1400/800	0.321 *	[72]
			CeO ₂	1350/850	0.05	[76]
			CeO ₂	1500/1000	0.22	[76]
			LaGa _{0.4} Co _{0.6} O _{3-δ}	1350/800	0.478	[31]
Three reactors chemical looping (M-TRCL)	ABO ₃	$ABO_3 \leftrightarrow ABO_{3-\delta} + 0.5\delta O_2$; $ABO_{3-\delta} + \delta H_2O \rightarrow ABO_3 + \delta H_2$	Sr _{0.4} La _{0.6} Mn _{0.6} Al _{0.4} O ₃	1350/1000	0.307	[75]
			Sr _{0.4} La _{0.6} Mn _{0.6} Al _{0.4} O _{3-δ}	1350/850	0.194	[76]
	Iron oxides (Fe ₂ O ₃)	$Fe_2O_3 + CH_4 \rightarrow FeO + 3CO_2 + 2H_2O$; $3FeO + H_2O \rightarrow Fe_3O_4 + H_2$; $2Fe_3O_4 + 0.5O_2 \rightarrow 3Fe_2O_3$	BaCe _{0.25} Mn _{0.75} O _{3-δ}	1350/850	0.140	[76]
			CaTi _{0.5} Mn _{0.5} O _{3-δ}	1350/1150	0.446	[78]
			K promoted-Fe ₂ O ₃ /Al ₂ O ₃	850/850	-	[97]
	Cerium oxide (Ce _x M _{1-x} O ₂)	$Ce_xM_{1-x}O_2 + \delta CH_4 \rightarrow Ce_xM_{1-x}O_{2-\delta} + \delta CO + 2\delta H_2$; $Ce_xM_{1-x}O_{2-\delta} + \delta H_2O \rightarrow Ce_xM_{1-x}O_2 + \delta H_2$	Fe ₂ O ₃ /La _{0.8} Sr _{0.2} FeO ₃	900/900	-	[98]
			Ce _{0.5} Fe _{0.5} O _{2-δ}	850/850	3.71	[87]
			5Ni/CeO ₂ -TiO ₂	900/900	2.098 *	[108]
Methane chemical looping process	CL-SMR		CeO ₂ -Fe ₂ O ₃ (Ce/Fe = 7/3)	850/700	1.116	[109]
			40 wt%			
			Ce-Fe-Zr-O/MgAl-LDO	850/850	1.0	[111]
	ABO ₃	$ABO_3 + \delta CH_4 \rightarrow ABO_{3-\delta} + \delta CO + 2\delta H_2$; $ABO_{3-\delta} + \delta H_2O \rightarrow ABO_3 + \delta H_2$	CeO ₂ -LaFeO ₃	800/800	3.38	[112]
			CeO ₂ /ZrO ₂	800/800	1.736	[114]
			LaCo _{0.6} Fe _{0.4} O ₃	700/700	2.22	[117]
			La _{0.5} Ce _{0.5} FeO ₃	925/925	0.6	[118]
			La _{0.85} MnFe _{0.15} O ₃	850/850	1.76	[122]
			Fe ₂ O ₃ /SiO ₂	900/900	~0.745 *	[135]
			Fe ₂ O ₃ /Al ₂ O ₃	900/900	~0.982 *	[135]
			Fe ₂ O ₃ /YSZ	900/900	~2.232 *	[135]
			Fe ₂ O ₃ /ZrO ₂	900/900	~2.977 *	[135]
			Fe ₂ O ₃ /MgAl ₂ O ₄	900/900	2.41~4.152 *	[135]
CL-WGS	Iron oxides (Fe ₂ O ₃)	$Fe_2O_3 + 3CO \rightarrow 2Fe + 3CO_2$; $3Fe + 4H_2O \rightarrow Fe_3O_4 + 4H_2$; $2Fe_3O_4 + 0.5O_2 \rightarrow 3Fe_2O_3$	Na-modified Fe ₂ O ₃ /Al ₂ O ₃	800/800	~13.3	[136]
			Fe ₂ O ₃ /ZrO ₂	850/850	~7.143 *	[137]
			Fe ₂ O ₃ /CeO ₂	850/850	~5.80 *	[137]
			Fe ₂ O ₃ /Ce _{0.6} Sm _{0.15} Zr _{0.25} O _{1.925}	850/850	2.422 *	[139]

Table 1. Cont.

Hydrogen Production Processes	OCs Type	Cyclic Reaction	OCs	Reaction Temperature (°C; Red/Ox)	H ₂ Yield (mmol/g)	Ref.
SCL process	Iron spinels (M _x Fe _{3-x} O ₄)	M _x Fe _{3-x} O ₄ + (2-x)H ₂ + CO → xMO + (3-x)Fe + CO ₂ + (3-x)H ₂ O; Fe + H ₂ O → Fe ₃ O ₄ + H ₂ ; xMO + (3-x)/3Fe ₃ O ₄ + 0.5xO ₂ → M _x Fe _{3-x} O ₄	3.3 wt% Al-modified NiFe ₂ O ₄	800/800	8.2	[143]
			ZnFeAlO _x	850/850	2.23	[144]
			CoFe ₂ O ₄	650/650	11.9	[145]
			Cu _{0.25} Co _{0.25} Fe _{2.5} O ₄	550/550	~11.9	[146]
	Brownmillerite (Ca ₂ Fe ₂ O ₅)	Ca ₂ Fe ₂ O ₅ + 3CO → 2CaO + 2Fe + 3CO ₂ ; 2CaO + 2Fe + 3H ₂ O → Ca ₂ Fe ₂ O ₅ + 3H ₂	Ca ₂ Fe ₂ O ₅	900/900	-	[149]
	Iron oxides (Fe ₂ O ₃)	Fe ₂ O ₃ + 2H ₂ + CO → 2Fe + CO ₂ + 2H ₂ O 3Fe + 4H ₂ O → Fe ₃ O ₄ + 4H ₂ ; 2Fe ₃ O ₄ + 0.5O ₂ → 3Fe ₂ O ₃	Fe-Ce-Ni based OCs (Ce/Ni/Fe ratio of 10/3/100) Fe ₂ O ₃	750/750 600/600	11.79 1.14	[158] [162]
		M _x Fe _{3-x} O ₄ + CO → xMO + (3-x)Fe + CO ₂ ; xMO + (3-x)FeO + H ₂ O → M _x Fe _{3-x} O ₄ + H ₂ ;	Co _{0.85} Fe _{2.15} O ₄ /m-ZrO ₂	600/600	3.944	[162]
	Iron spinels (M _x Fe _{3-x} O ₄)	2TiO ₂ → Ti ₂ O ₃ + 0.5O ₂ ; Ti ₂ O ₃ + H ₂ O → 2TiO ₂ + H ₂ (hv + TiO ₂ → e ⁻ + h ⁺ ; e ⁻ + Ti ⁴⁺ → Ti ³⁺ ; h ⁺ + O ₂ → O ₂ + Vo; Vo + H ₂ O → Vo(H ₂ O) Ti ³⁺ -Vo (H ₂ O)-Ti ³⁺ → Ti ⁴⁺ -O-Ti ⁴⁺ + H ₂)	TiO ₂	room temperature/500-600	0.0188 *	[34]
			0.5 wt% Fe/TiO ₂	room temperature/600	0.0334	[163]
			TiO ₂	room temperature/350	0.75 × 10 ⁻³	[165]
			Fe/TiO ₂	room temperature/350	9.73 × 10 ⁻³	[165]
			Cu/TiO ₂	room temperature/350	7.13 × 10 ⁻³	[165]
			Ni-promoted Cu/TiO ₂	room temperature/350	0.027	[165]
			0.5% Ni-promoted Cu/TiO ₂	room temperature/350	0.0306	[166]

* The unit of raw hydrogen production data is “mL/g”, which can be converted into “mmol/g” by the formula PV = nRT with hypothesis of STP condition.

For the two-step TCWS cycle, various OCs, such as iron oxides, zinc oxides, cerium oxides, and perovskite have been widely studied. Among them, cerium oxides have attracted particular attention due to their high structural stability and water-splitting conversion. However, the relatively low reduction degree during the redox cycle, rendering a low hydrogen yield (0.72~7.58 mL/g), greatly hampered its practical applications. Recent work showed that perovskite oxides are promising candidates for two-step TCWS with hydrogen yields of up to 3.13 to 10.71 mL/g, since their redox properties can be facilely modulated by tuning the A/B sites. This gives a clue that constructing composite materials to adjust the redox potential suitable for oxygen desorption and water-splitting should be the key for improving the hydrogen productivity. Furthermore, the sintering problems, thermal shocks, and long single redox cycle time, induced by application of high thermal reduction temperature and low water-splitting temperature, represents another huge obstacle for commercialization of this process. Therefore, introducing support to stabilize the OCs and exploring isothermal redox reactions would be effective in improving the process efficiency for hydrogen production.

Compared to the two-step TCWS cycle, introducing reducing gas, such as methane, carbon monoxide, and syngas, to reduce the oxygen carrier is capable of notably decreasing the reaction temperature to below 1000 °C while enhancing the available oxygen capacity, which significantly decreases the energy consumption, slows down the sintering of OCs, and improve the yield of hydrogen to 13.44~267.63 mL/g. As for methane-driven reduction, valuable syngas with H₂/CO ratio of two for Fischer–Tropsch synthesis and methanol production is produced when a suitable OC is selected. Upon OCs with high reducibility applied, the reducing gaseous can be totally combusted with generation of high concentration CO₂ (and H₂O). All these processes greatly inhibit the side reactions and reduce the burden for gas separation and CO₂ caption, rendering improved efficiency and lowered cost for hydrogen production. Among the investigated OCs, iron-based oxides are among the most studied materials due to the virtues of low-cost, environmentally benign features with the high capacity to donate lattice oxygen by varying the valence state of Fe cations. Herein, the relatively low reactivity of iron cations towards conversion of these reducing agents and general trade-off between selectivity and hydrogen yield represent the main challenge for widespread application of these techniques. For CL-SMR with the aim of methane partial oxidation, the reactivity for C–H bond cleavage can be improved by introducing reactive promoters (e.g., Ni and Cu), creating oxygen vacancies, and engineering the morphology of OCs to provide more active centers. To break the trade-off between selectivity and hydrogen productivity, the key lies in a balance between oxygen supply and methane decomposition, which sustainably restrains the coke formation and increases the oxygen capacity for water-splitting. Based on available results, constructing OCs with core-shell structure, composite oxides, and poisoning active Fe⁰ by forming an alloy or covering with oxides are effective methods to inhibit the carbon deposition. When total combustion of reducing agents (processes of methane-driven three step CLWS, CL-WGS, and SCL) is desired, the reduction degree of OC and hydrogen productivity can be improved by introducing MIEC support, alkali promoters, and some active metals (e.g., Co and Cu), which contribute to promote oxygen mobility and oxygen capacity.

In the case of PTC cycle, the reduction of OCs was realized at ambient temperature with hydrogen production at low temperature (140~600 °C), which greatly suppressed the sintering of OCs. More importantly, the solar energy can be well converted to chemical energy, implementing production of green hydrogen. Up to now, the studied OCs are still limited to bare or modified titanium oxide. Although much progress has been achieved in restraining the recombination rate of EHPs and enhancing the reactivity towards water-splitting, which resulted in improved redox performance, the hydrogen yield (0.0168~0.685 mL/g) is still lower than the above CLWS process, because the photo-driven reduction mainly drives the surface lattice oxygen desorption of OCs, and the amount of desorbed oxygen desorption is limited. In addition to exploring advanced OCs that release more oxygen in the photothermal reduction process, it is anticipated that a combination

of suitable reducing gas and photothermal reduction would be promising to significantly enhance the efficiency of hydrogen production.

Over the past 30 years, great effort has been made to develop various OCs by engineering the composition of oxides, including simple metal oxides (cerium oxides, iron oxides, titanium oxides, etc.), composite metal oxides (e.g., perovskite, spinel and garnets), and mixed oxides (e.g., supported OCs, $\text{Fe}_2\text{O}_3\text{-CeO}_2$, and $\text{CeO}_2\text{-TiO}_2$), which greatly improved the redox performance in CLWS processes [31,48,58,97,104,108,110,125,145,167]. In addition, modulating the microstructure during the redox reactions also exerts a big influence on the performance due to the modulating of surface active centers, concentration of oxygen vacancies, or metal–oxygen bond strength. For example, Li et al. [124] prepared a kind of $\text{Fe}_2\text{O}_3@\text{La}_x\text{Sr}_{1-x}\text{FeO}_{3-\delta}$ (LSF) OC with core-shell structure and showed that the LSF shell with high methane-to-syngas selectivity could reduce the contact between methane and Fe_2O_3 , which enables a high CO selectivity above 98%, while the Fe_2O_3 core can donate oxygen to the LSF shell that greatly enhances the oxygen capacity of this OC. Qian et al. [78] found that the crystal phase of $\text{CaTi}_{0.5}\text{Mn}_{0.5}\text{O}_{3-\delta}$ OCs could change from an orthogonal phase to a cubic phase during the thermal reduction process, which greatly promoted the desorption of lattice oxygen. Furthermore, the OCs with cubic phase integrate the virtues of high entropy and medium enthalpy, which improves the reactivity for hydrogen production via water-splitting. Wang et al. [121] demonstrated that a specific Fe^0 @ oxide mediate with core-shell structure was formed when $\text{La}_{0.6}\text{Sr}_{0.4}\text{Fe}_{0.8}\text{Al}_{0.2}\text{O}_{3-\delta}$ was reduced by methane, which notably reduced the direct contact between methane and generated Fe^0 species, restrained the coke deposition, and improved the syngas productivity. Cui et al. [144] indicated that the active component (Fe) of ZnFeAlO_x OCs reversibly exsolved and dissolved from the interface structure of Zn-Fe-Al-O mixed spinel during a CL-WGS process, which can effectively hinder the migration and agglomeration of iron ions and avoid sintering of materials. These results clearly indicate that adjusting the structure changes during redox reactions represents an effective method to modulate the redox performance for CLWS processes.

At present, selection of OCs for CLWS reactions mainly relies on screening method. This is mainly ascribed to the harsh reaction conditions and dynamic structural evolution during redox reactions, which poses a huge challenge for comprehensively understanding the reaction mechanism and designing advanced OCs for CLWS reactions. Future studies should pay more attention to establish a more precise structure–function relationship with the help of in situ characterization, theoretical calculations, and thermodynamic analysis to provide a theoretical basis and development direction for the design of new efficient long-life OCs. Furthermore, according to the pioneering studies, the research focus of OCs is gradually transferred from simple metal oxides to composite oxides (e.g., perovskite) and mixed oxides due to the feasibility of modulating the redox properties by altering the composition of OCs or synergy between different oxides, which bypasses the shortcomings of single metal oxides, and improves the performance of hydrogen production. Therefore, exploring composite oxides to precisely control the metal–oxygen bond strength and mixed oxides to integrate the advantages of different oxides would be an effective strategy for further improving the redox performance of OCs.

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