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## Chemical looping beyond combustion – a perspective

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As a promising approach for carbon dioxide capture, chemical looping combustion has been extensively investigated for more than two decades. However, the chemical looping strategy can be and has been extended well beyond carbon capture. In fact, significant impacts on emission reduction, energy conservation, and value-creation can be anticipated from chemical looping beyond combustion (CLBC). This article aims to demonstrate the versatility and transformational benefits of CLBC. Specifically, we focus on the use of oxygen carriers or redox catalysts for chemical production – a \$4 trillion industry that consumes 40.9 quadrillion BTU of energy. Compared to state-of-the-art chemical production technologies, we illustrate that chemical looping offers significant opportunities for process intensification and exergy loss minimization. In many cases, an order of magnitude reduction in energy consumption and CO<sub>2</sub> emission can be realized without the needs for carbon dioxide capture. In addition to providing various CLBC examples, this article elaborates on generalized design principles for CLBC, potential benefits and pitfalls, as well as redox catalyst selection, design, optimization, and redox reaction mechanism.

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### Broader context

The chemical industry consumes more than 40 quadrillion BTU of energy while emitting 2.5 Gigatons of carbon dioxide each year. Meanwhile, many established chemical processes are extensively optimized, providing limited space for further energy savings and emission reductions. Chemical looping offers exciting new opportunities to the aforementioned challenges through process intensification. Facilitated by a redox catalyst, a rationally designed chemical looping scheme can reduce the number of unit operations and significantly decrease the exergy loss for chemical production. In some cases, the chemical looping strategy can facilitate a chemical conversion that would be otherwise infeasible in a conventional reaction scheme. While a number of chemical looping based chemical production approaches have shown promising results, this concept can be further expanded to a significantly wider sets of products using a variety of feedstock and energy sources. Given the exciting opportunities offered by chemical looping beyond combustion (CLBC) and the emissions and energy demand in the current chemical production practice, the CLBC design principles and redox catalyst selection/optimization strategies covered in this article can provide useful information for researchers in the general areas of chemical looping, catalysis, particle technology, and process intensification.

## 1. Introduction

Although the term “chemical looping” was first minted by Richter and Knoche in 1983 in the context of reducing exergy loss in fossil fuel combustion,<sup>1</sup> the concept of chemical looping (CL),

*i.e.* decomposition of a chemical reaction into multiple sub-reactions facilitated by solid reaction intermediates, was investigated long before that.<sup>2,3</sup> Owing to the pressing demand for carbon emissions reduction,<sup>4,5</sup> chemical looping combustion (CLC) has been studied extensively over the past three decades as a new technology for power generation with integrated CO<sub>2</sub> capture,<sup>6–22</sup> as evidenced by more than 2400 peer-reviewed publications to date (based on the chemical abstracts service). In comparison, research related to chemical looping strategies prior to 1983 was scattered and often dealt with concepts other than power generation and CO<sub>2</sub> capture.<sup>2,23–25</sup> A good example is the commercially implemented steam-iron process for hydrogen production, which was later replaced by methane reforming in the 1930s.<sup>23</sup> However, besides the limitations from the lack of global policies on fighting global warming *via* carbon capture and storage (CCS)-related technologies, the lack of fundamental

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understanding and absence of effective strategies to design and optimize the reaction intermediates, also known as (oxygen) carriers, have largely hindered the research progress in the “pre-CLC” era. We note that the extensive investigation of the CLC technology<sup>26–37</sup> has built up significant background knowledge and fundamental insights for potential breakthroughs in chemical looping applications well beyond combustion and power generation.<sup>24,25,38–51</sup> It is, therefore, not too surprising to see that the extension of the chemical looping strategy has drawn significant attention over the past decade. In fact, journal articles covering such topics increased by nearly 10-fold over this period.

Although there have been quite a few excellent reviews and books covering the topic of chemical looping, they tend to focus on CLC<sup>26–35</sup> and, in some cases, chemical looping reforming (CLR) both with and without being integrated with CO<sub>2</sub> or H<sub>2</sub>O splitting.<sup>38–53</sup> A recent review by Zeng *et al.*<sup>21</sup> provided comprehensive discussions of the various aspects of chemical looping including chemical looping for the production of chemicals. However, CLC and CLR related topics were still the primary focus, and many chemical production cases, as summarized in Table 1, have not been covered. Another review by Bayham *et al.*<sup>22</sup> gave an overview of CLC for power production with CO<sub>2</sub> capture and CLR for hydrogen or syngas production with carbon capture. It mainly focused on the potential process configurations (reaction systems and reactor design) and applications. As a highly versatile and effective process intensification strategy, the chemical looping concept can be applied to many applications beyond CO<sub>2</sub> capture and reforming with significant merits in efficiency improvements, emission reduction, and potential cost savings. This article aims to offer a perspective on various potential applications of chemical looping beyond combustion (CLBC) and the generalized design principles for oxygen carrier or redox catalysts in chemical looping processes.

## 2. The case for chemical looping beyond combustion

### 2.1. Chemical looping – the general principle

As illustrated in Table 1, chemical looping can take various forms for numerous applications. Generalized schematics and representative reactions of different chemical looping processes are summarized in Fig. 1 and Table 1, respectively. As can be seen, reactive separation facilitated by solid (oxygen, nitrogen, and hydrogen) carrier particles represents an important underlying feature of all chemical looping reactions and processes. In chemical looping processes, oxygen carrier, usually composed of metal oxides, acts as an oxygen reservoir for oxygen donation and regeneration.<sup>21</sup> Nitrogen carrier composed of nitride or nitride-hydride mixed compounds enables the storage and release of N or/and H in the redox cycle for chemical looping ammonia synthesis.<sup>98</sup> In a similar manner, hydride based hydrogen carrier allows the storage and release of hydrogen to participate in in chemical reactions or shift the reaction equilibrium.<sup>106</sup> Although recent research on nitrogen,<sup>98–105,107,108</sup> and hydrogen<sup>106</sup> carriers

offer significant new opportunities, the vast majority of the carriers investigated to date are oxygen carrying agents. In order to maintain coherency, we elected to focus exclusively on chemical looping approaches for chemical production using oxygen carriers, also known as redox catalysts. Table 1 summarizes both the key CLBC schemes reported to date and promising CLBC schemes for future investigations. To further illustrate the thermodynamic favorability of the various potential reaction pathways, Fig. 2 summarizes potential reaction pathways and Gibbs free energy changes ( $\Delta G$ ) among various carbon-containing feedstock and products. To maintain consistency, 1 bar oxygen partial pressure ( $P_{O_2}$ ) was assumed in all cases. Fig. 2 is intended to be used as a screen tool to determine the thermodynamically favorable reactants, products, and CLBC schemes.<sup>49</sup> It is also important to note that while one can anticipate similar trends in relative ease of product formation irrespective to the oxygen carrier used in CLBC, the equilibrium  $P_{O_2}$  of oxygen carriers, which is dependent upon both the composition/phase of the redox pair and the reaction temperature, can span more than 20 orders of magnitudes. This large degree of freedom in  $P_{O_2}$  and the ability of chemical looping to decouple a single overall reaction into multiple redox steps, provides unique flexibility in “manipulating” the extent and feasibility of certain chemical reactions, as will be illustrated in Section 2.2. Besides redox based CLBC schemes, this article also covers the use of CO<sub>2</sub> sorbents (*i.e.* calcium looping) to enhance the oxygen carrier-based chemical looping approaches, as will be elaborated in Section 4.4.

A typical chemical looping process involving oxygen-carrying agents is composed of two or more reduction and oxidation steps that form a redox loop. In its simplest form, an oxygen carrier or redox catalyst first donates its lattice oxygen under a low oxygen partial pressure environment. The reduced oxygen carrier is subsequently exposed to an oxidant for the replenishment of its lattice oxygen, thereby completing a two-step redox loop. As illustrated in Table 1, the use of oxygen carrier offers significant flexibility and opportunity for process intensification by breaking an overall reaction into sub-reactions because: (i) besides acting as a reactant, the oxygen carrier also acts as a mass separation agent since the looping reactions prevent the mixing among the products from each sub-step; (ii) the carrier's oxygen donation properties can be tailored for improved thermal management of the overall process; and (iii) the reduction or oxidation steps can be further broken into multiple sub-steps.<sup>15,21</sup> When designed properly, an oxygen carrier can be compatible with various oxidizing and reducing agents to facilitate the generation of multiple value-added products with minimal separation requirements.

From a thermodynamic standpoint, the oxygen donation ability of oxygen carriers/redox catalysts can be evaluated by the equilibrium oxygen partial pressure ( $P_{O_2}$ ) of the reaction  $MeO_x \rightarrow MeO_{x-1} + 1/2O_2$ .<sup>49</sup> Redox pairs with high  $P_{O_2}$ s, which are suitable for chemical looping air separation (CLAS), chemical looping with oxygen uncoupling (CLOU), or CLC, can only be regenerated with air. As applications summarized in Table 1, CLBC can take advantage of redox pairs with a significantly wider range of  $P_{O_2}$ s. As such, valuable products can be produced



Table 1 Representative redox schemes for chemical looping beyond combustion. The redox reactions are named from (R1) to (R41)

Feedstock	Products	Product generation	Balance of the loop	Name of the process <sup>a</sup>	Sample oxygen/nitrogen carriers
Air	Air separation	(R1) $\text{MeO}_x \rightarrow \text{MeO}_{x-1} + 1/2\text{O}_2$	(R2) $\text{MeO}_{x-1} + 1/2\text{O}_2 \rightarrow \text{MeO}_x$	CLAS	$\text{CuO}$ , <sup>49</sup> perovskite <sup>54,55</sup>
Methane	Synthesis gas	(R3) $\text{CH}_4 + \text{MeO}_x \rightarrow \text{MeO}_{x-1} + \text{CO} + 2\text{H}_2$	(R4) $\text{MeO}_{x-1} + 1/2\text{O}_2 \rightarrow \text{MeO}_x$ (R5) $\text{MeO}_{x-1} + \text{H}_2\text{O} \rightarrow \text{MeO}_x + \text{H}_2$ (R6) $\text{MeO}_{x-1} + \text{CO}_2 \rightarrow \text{MeO}_x + \text{CO}$	CLR	$\text{NiO}$ , <sup>56,57</sup> $\text{FeO}_x$ , <sup>58-62</sup> $\text{CeO}_2$ , <sup>63-65</sup> perovskite, <sup>66-68</sup> hexaaluminate <sup>69,70</sup>
	Ethylene	(R7) $\text{CH}_4 + \text{MeO}_x \rightarrow \text{MeO}_{x-1} + 1/2\text{C}_2\text{H}_4 + \text{H}_2\text{O}$	(R8) $\text{MeO}_{x-1} + 1/2\text{O}_2 \rightarrow \text{MeO}_x$	CL-OCM	Alkali modified Mn, Fe oxides <sup>71,72</sup>
	Benzene	(R9) $2/3\text{CH}_4 + \text{MeO}_x \rightarrow \text{MeO}_{x-1} + 1/9\text{C}_6\text{H}_6 + \text{H}_2\text{O}$	(R10) $\text{MeO}_{x-1} + 1/2\text{O}_2 \rightarrow \text{MeO}_x$	CL-DHA	Mo/Zn ZSM-5 + Fe oxide <sup>73,74</sup>
	Methanol	(R11) $\text{CH}_4 + \text{MeO}_x \rightarrow \text{MeO}_{x-1} + \text{CH}_3\text{OH}$	(R12) $\text{MeO}_{x-1} + \text{H}_2\text{O} \rightarrow \text{MeO}_x + \text{H}_2$	CL-Selective oxidation	Copper-exchanged zeolites <sup>75,76</sup>
Methanol	Formaldehyde	(R13) $\text{CH}_3\text{OH} + \text{MeO}_x \rightarrow \text{MeO}_{x-1} + \text{CH}_2\text{O} + \text{H}_2\text{O}$	(R14) $\text{MeO}_{x-1} + 1/2\text{O}_2 \rightarrow \text{MeO}_x$	CL-Selective oxidation	$\text{FeMoO}_4\text{-Fe}_2\text{O}_3$ <sup>77</sup>
$\text{C}_{2+}$ hydrocarbon	Ethylene oxide	(R15) $\text{C}_2\text{H}_4 + \text{MeO}_x \rightarrow \text{MeO}_{x-1} + \text{C}_2\text{H}_4\text{O}$	(R19) $\text{MeO}_{x-1} + 1/2\text{O}_2 \rightarrow \text{MeO}_x$ or (R21) $\text{MeO}_{x-1} + \text{CO}_2 \rightarrow \text{MeO}_x + \text{CO}$	CL-Epoxidation	$\text{Ag/SrFeO}_3$ <sup>78</sup>
	Ethylene	(R16) $\text{C}_2\text{H}_6 + \text{MeO}_x \rightarrow \text{MeO}_{x-1} + \text{C}_2\text{H}_4 + \text{H}_2\text{O}$		CL-ODH	Alkali modified Mn and Fe oxides <sup>79-83</sup>
	Propylene	(R17) $\text{C}_3\text{H}_8 + \text{MeO}_x \rightarrow \text{MeO}_{x-1} + \text{C}_3\text{H}_6 + \text{H}_2\text{O}$		CL-ODH	Mn and La perovskite, <sup>84</sup> and $\text{VO}_x$ <sup>85</sup>
	Propionaldehyde	(R18) $1/2\text{C}_3\text{H}_8 + \text{MeO}_x \rightarrow \text{MeO}_{x-1} + 1/2\text{C}_3\text{H}_6\text{O} + 1/2\text{H}_2\text{O}$		CL-Selective oxidation	— <sup>b</sup>
	Maleic anhydride	(R20) $1/7\text{C}_4\text{H}_{10} + \text{MeO}_x \rightarrow \text{MeO}_{x-1} + 1/7\text{C}_4\text{H}_2\text{O}_3 + 4/7\text{H}_2\text{O}$		CL-Selective oxidation	Vanadium phosphorous oxygen (VPO) <sup>86</sup>
	Butadiene	(R22) $\text{C}_4\text{H}_8 + \text{MeO}_x \rightarrow \text{MeO}_{x-1} + \text{C}_4\text{H}_6 + \text{H}_2\text{O}$		CL-ODH	—
	Ethylene	(R23) $\text{C}_6\text{H}_{14} + \text{MeO}_x \rightarrow \text{MeO}_{x-1} + 3\text{C}_2\text{H}_4 + \text{H}_2\text{O}$		CL-Oxidative cracking	Alkali modified Mn and Fe oxides <sup>87</sup>
Aromatic	Styrene	(R24) $\text{C}_8\text{H}_{10} + \text{MeO}_x \rightarrow \text{MeO}_{x-1} + \text{C}_8\text{H}_8 + \text{H}_2\text{O}$	(R26) $\text{MeO}_{x-1} + 1/2\text{O}_2 \rightarrow \text{MeO}_x$	CL-ODH	—
	Methylstyrene	(R25) $\text{C}_9\text{H}_{12} + \text{MeO}_x \rightarrow \text{MeO}_{x-1} + \text{C}_9\text{H}_{10} + \text{H}_2\text{O}$			—
Biomass	Synthesis gas	(R27) $\text{C}_x\text{H}_y\text{O}_z + \text{MeO}_x \rightarrow \text{MeO}_{x-1} + \text{synthesis gas}$	(R28) $\text{MeO}_{x-1} + 1/2\text{O}_2 \rightarrow \text{MeO}_x$	CLR	Fe oxide <sup>44</sup>
$\text{H}_2\text{O}$	Hydrogen	(R29) $\text{H}_2\text{O} + \text{MeO}_{x-1} \rightarrow \text{MeO}_x + \text{H}_2$	(R30) $\text{MeO}_x \rightarrow \text{MeO}_{x-1} + 1/2\text{O}_2$	Thermochemical splitting	Perovskite, <sup>88</sup> $\text{CeO}_2$ , <sup>64</sup> Fe oxide <sup>89</sup>
			(R31) $\text{MeO}_x + \text{CO} \rightarrow \text{CO}_2 + \text{MeO}_{x-1}$	CL-WGS	Fe oxide <sup>90-92</sup>
$\text{CO}_2$	CO	(R32) $\text{CO}_2 + \text{MeO}_{x-1} \rightarrow \text{MeO}_x + \text{CO}$	(R33) $\text{MeO}_x \rightarrow \text{MeO}_{x-1} + 1/2\text{O}_2$	Thermochemical splitting	Perovskite, <sup>67</sup> $\text{CeO}_2$ , <sup>93</sup> Fe oxide <sup>94</sup>
			(R34) $\text{MeO}_x + \text{H}_2 \rightarrow \text{H}_2\text{O} + \text{MeO}_{x-1}$	CL-RWGS	Perovskite, <sup>95</sup> Fe oxide <sup>96,97</sup>
$\text{N}_2/\text{H}_2$	$\text{NH}_3$	(R35) $3/2\text{H}_2 + \text{MeN}_x \rightarrow \text{MeN}_{x-1} + \text{NH}_3$ (R37) $\text{H}_2\text{O} + \text{MeN}_x \rightarrow \text{MeO} + x\text{NH}_3 + (1 - 3x/2)\text{H}_2$ (R40) $3/2\text{H}_2 + 1/\beta\text{MeH}_x\text{N}_\beta \rightarrow 1/\beta\text{MeH}_x + \text{NH}_3$	(R36) $1/2\text{N}_2 + \text{MeN}_{x-1} \rightarrow \text{MeN}_x$ (R38) $\text{MeO} + [2/(4m + n)]\text{C}_m\text{H}_n \rightarrow \text{Me} + [n/(4m + n)]\text{H}_2\text{O} + [2m/(4m + n)]\text{CO}_2$ (R39) $x/2\text{N}_2 + \text{Me} \rightarrow \text{MeN}_x$ (R41) $1/\beta\text{MeH}_x + 1/2\text{N}_2 \rightarrow 1/\beta\text{MeH}_x\text{N}_\beta$	CL-ammonia synthesis	Transition metal nitride <sup>98</sup>
					Metal nitride/metal oxide/metal <sup>99-101</sup>
					Nitrogen-containing hydride/hydride <sup>98-105</sup>

<sup>a</sup> CLAS, chemical looping air separation; CL-OCM, chemical looping oxidative coupling of methane; CL-DHA, chemical looping dehydroaromatization; CL-ODH, chemical looping oxidative dehydrogenation; CL-WGS, chemical looping water-gas shift; CL-RWGS, chemical looping reverse water-gas shift. <sup>b</sup> Exemplary, promising CLBC schemes that has yet to be investigated.

in both the oxygen carrier reduction and re-oxidation steps. In the reduction or oxygen donation step, CLBC can valorize light alkanes, aromatics or biomass into chemicals *via* chemical looping partial oxidation (CLPOx). Although  $\text{CO}_2$  and  $\text{H}_2\text{O}$  are the most stable forms of the oxidation products (other than carbonates), many chemical products, including hydrogen, synthesis gas, alkene, aromatic, alcohol, aldehyde, epoxide, or maleic anhydride, are thermodynamically feasible as illustrated

in Fig. 2b. The key challenge for high yields towards these partial oxidation products resides in the design of selective redox catalysts to minimize  $\text{CO}_x$  and other byproduct formation, as will be elaborated in Section 3.2. In the regeneration step, hydrogen can be produced *via* water splitting or CO from  $\text{CO}_2$  splitting. Selective hydrogenation reactions can also be carried out during the re-oxidation of tailored redox catalysts. Although the chemical looping cycles can be more complex





Fig. 1 Schematic illustration of the chemical looping strategy and its potential applications.



Fig. 2 Normalized reaction Gibbs free energy change ( $\Delta G$ ) per mole of carbon at 700 °C for various potential chemical products from C1–C9 hydrocarbons. Bars with different colors represent the  $\Delta G$  for the partial oxidation of hydrocarbons to the corresponding products. For consistency, 1 atm gaseous oxygen is assumed to be the oxidant. To calculate  $\Delta G$  with a specific redox pair as the oxidant (e.g.  $\text{Mn}_2\text{O}_4/\text{MnO}$ ), one can use the corresponding equilibrium oxygen partial pressure ( $P_{\text{O}_2}$ ) of the redox pair at the temperature to account for the oxygen chemical potential change relative to 1 atm  $\text{O}_2$ .

than conventional processes in the presence of heterogeneous catalysts, they do offer multiple potential advantages from both product selectivity and process efficiency viewpoints, as will be further illustrated in the following sections.

## 2.2. The case for chemical looping beyond combustion – a second law perspective

As discussed in Section 2.1, a common feature of chemical looping is the ability to facilitate chemical reactions with *in situ* product separation. With separation being the most energy-intensive step in the chemical industry, reactive separation enabled by chemical looping offers excellent opportunities for energy savings and emission reductions. The potential benefits of chemical looping, however, goes well-beyond the simplification of product separations. With strategically designed reaction scheme and intermediates, chemical looping can be intrinsically advantageous over conventional approaches from a second law of

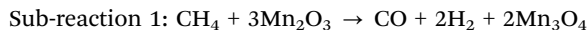
thermodynamics perspective. The potential second law (exergy) savings *via* chemical looping result from the following aspects:

**a. Exergy savings *via* reactive separation.** From a separation standpoint, the redox catalyst in chemical looping acts as a (reactive) mass separation agent, which facilitates feedstock and/or product separation. As such, the driving force ( $\Delta G$ ) for the overall chemical reaction, which is broken down into two or more sub-reactions in chemical looping, can be utilized for separation in an integrated manner. This represents a distinct advantage over conventional separation, in which chemical potential gradients among the various components in a mixture are created *via* extensive cooling or heating (e.g. distillation, cryogenic distillation, absorption-stripping, *etc.*) and/or compression (e.g. membrane, adsorption, and cryogenic distillation). With tunable thermodynamic properties, the driving forces for sub-reactions (and separation) can also be adjusted in chemical looping to optimize exergy savings. This is exemplified by the partial oxidation (POx) of methane: Under the conventional scheme, synthesis gas is produced from methane in two major steps, that is, cryogenic air separation and methane partial oxidation. Air separation is a thermodynamically uphill process requiring a minimum work input of  $6.2 \text{ kJ mol}^{-1} \text{ O}_2$  for a hypothetical and fully reversible separation process. The actual energy consumption for commercial cryogenic air separation, which requires extensive gas compression, liquefaction, and distillation, is approximately  $25 \text{ kJ mol}^{-1} \text{ O}_2$  (at 1 bar). This corresponds to an exergetic efficiency of  $\sim 25\%$ .<sup>109</sup> Even under an idealized scenario,<sup>110</sup> conventional air separation would still lead to an exergy loss of  $5.3 \text{ kJ mol}^{-1} \text{ O}_2$ , resulting in a total energy consumption of  $11.5 \text{ kJ mol}^{-1} \text{ O}_2$ . As such, conventional methane POx, which consumes gaseous  $\text{O}_2$  at a  $\text{CH}_4:\text{O}_2$  stoichiometric ratio of 2 : 1, would consume at least 5.8 kJ of energy in air separation alone for each mole of methane converted. Methane POx, on the other hand, is thermodynamically highly favored with  $\Delta G^0$  of  $-86.7 \text{ kJ mol}^{-1}$  of methane converted. Chemical looping POx can take advantage of the large thermodynamic driving force for methane POx and integrate it for air

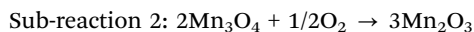




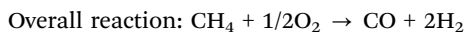
separation. Using  $\text{Mn}_2\text{O}_3$  as an example, the two chemical looping sub-reactions are each favored thermodynamically:



$$\Delta G_1^0 = -9.8 \text{ kJ mol}^{-1}$$



$$\Delta G_2^0 = -76.9 \text{ kJ mol}^{-1}$$



$$\Delta G^0 = -86.7 \text{ kJ mol}^{-1}$$

As can be seen, the thermodynamic driving force required for air separation is embedded in the chemical looping reactions, thereby eliminating the associated energy consumptions and exergy loss. It is further noted that, even when the chemical looping scheme is used for air separation only, it can be significantly less energy-intensive than cryogenic air separation.<sup>54,55</sup>

*b. Exergy savings via in situ conversion of byproducts.* The production of value-added chemicals can be intensified by chemical looping, *via* selectively oxidizing the reaction byproducts. As exemplified in Table 1, this principle has been applied to olefin production *via* oxidative coupling of methane, oxidative dehydrogenation, and naphtha oxidative cracking. From a second law standpoint, chemical looping can lead to exergy savings in the following two ways. On one hand, the *in situ* oxidation of byproducts such as hydrogen can compensate for the thermal energy required by endothermic reactions, *e.g.* the non-oxidative coupling of methane or ethane cracking/dehydrogenation. Compared to conventional approaches where hydrocarbon fuels are combusted to neutralize the reaction endothermicity *via* indirect heat transfer through cracker furnaces, *in situ* combustion of hydrogen is intrinsically advantageous due to its lower exergy rate, *i.e.* the ratio between the exergy and enthalpy of a fuel compared to that of hydrocarbons. Fig. 3 illustrates the exergy loss for ideal steam cracking and chemical looping oxidative dehydrogenation (CL-ODH) steps,<sup>111</sup> assuming no heat loss in neither case. When practical factors, such as the temperature difference for indirect heat transfer, are considered, a higher exergy loss can be anticipated for conventional cracking.

A second aspect of exergy savings *via* byproduct oxidation is through enhanced product yields. For instance, steam cracking

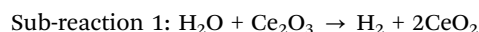
of both ethane and naphtha are equilibrium-limited.<sup>112</sup> *In situ* combustion of the hydrogen byproduct significantly increases the olefin yield from 50% to nearly 70% in a lab-scale reactor at 850 °C while decreasing the volumetric flow rate of non-condensable product gases by ~40%.<sup>81</sup> The corresponding energy savings in product compression and separation alone correspond to 554  $\text{kJ}_{\text{th}} \text{mol}^{-1}$  of ethylene produced.<sup>113</sup> Due to these reasons, process analyses indicate that the chemical looping approach can lead to more than 80% energy savings for ethane conversion and 50% energy savings for naphtha cracking.<sup>112,113</sup>

*c. Ability to circumvent second law limitations by the redox loop.*

Another potential advantage of dissecting an overall reaction into sub-reactions resides in a higher degree of freedom in terms of thermodynamic characteristics for the individual sub-reactions. Moreover, the operating conditions for each sub-step, *e.g.* temperature and pressure, can be adjusted independently to maximize the product yields from an equilibrium standpoint. A good example of using the chemical looping strategy to circumvent second law limitations of an overall reaction is thermochemical water-splitting to produce hydrogen:



Spontaneous decomposition of water (equilibrium constant,  $K \geq 1$ ) only occurs at ~4100 °C or higher, making thermochemical water-splitting far from practical. This is mainly due to the high reaction enthalpy ( $\Delta H^0 = 241.5 \text{ kJ mol}^{-1}$ ) and the relatively small reaction entropy ( $\Delta S^0 = 43.5 \text{ J K}^{-1}$ ). Using the chemical looping principle, however, the overall reaction is broken down into two sub-reactions, as shown in Fig. 4. Take a ceria-based oxygen carrier as an example:



$$\Delta H_1^0 = -126.6 \text{ kJ mol}^{-1} \quad \Delta S_1^0 = -82.3 \text{ J K}^{-1} \quad \Delta G_1^0 = -104.1 \text{ kJ mol}^{-1}$$



$$\Delta H_2^0 = 368.2 \text{ kJ mol}^{-1} \quad \Delta S_2^0 = 125.9 \text{ J K}^{-1} \quad \Delta G_2^0 = 333.7 \text{ kJ mol}^{-1}$$



$$\Delta H^0 = 241.5 \text{ kJ mol}^{-1} \quad \Delta S^0 = 43.5 \text{ J K}^{-1} \quad \Delta G^0 = 229.6 \text{ kJ mol}^{-1}$$

Using ceria as the oxygen carrier, the exothermic water-splitting reaction (sub-reaction 1) is favored at lower temperatures<sup>114</sup> but

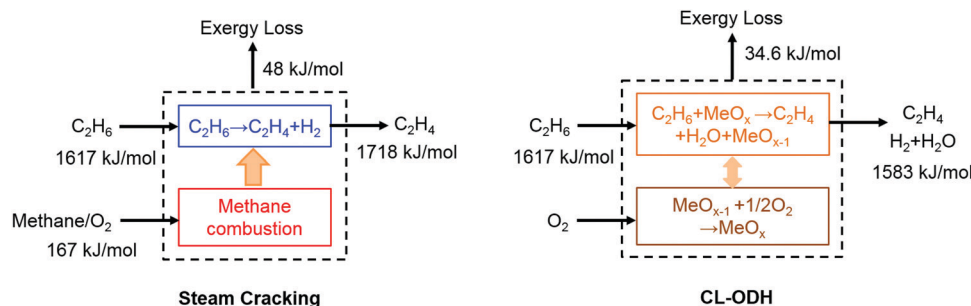


Fig. 3 Availability analysis and exergy loss for idealized steam cracking and chemical looping-oxidative dehydrogenation (CL-ODH) schemes.



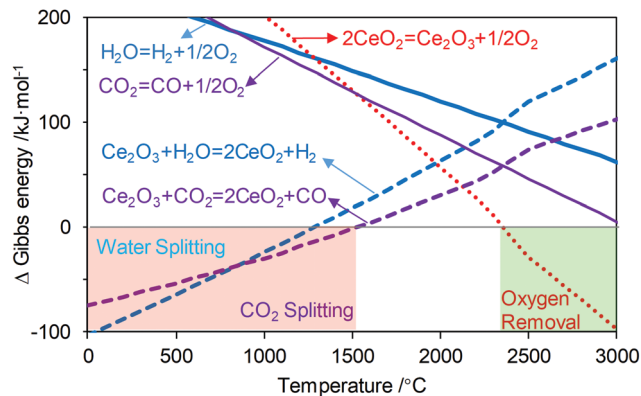


Fig. 4 Ellingham diagram of thermal and chemical looping water and CO<sub>2</sub> splitting. Light green and red regions represent thermodynamically spontaneous ( $K \geq 1$ ) reduction and oxidation reactions, respectively. In practice, a reaction would occur when  $K < 1$ .

remains favorable up to 1300 °C ( $K_{\text{eq}} \approx 1$ ). While the endothermic CeO<sub>2</sub> decomposition reaction is equilibrium-limited at low temperatures, the large entropy term, which increases with temperature, leads to spontaneous decomposition at 2300 °C, or 1800 °C lower than the direct water-splitting temperature ( $P_{\text{O}_2} = 1$  bar). While one can argue that 2300 °C is still too high, CeO<sub>2</sub> can be decomposed at much lower temperatures ( $\sim 1200$  °C) in practice by: (i) creating a low  $P_{\text{O}_2}$  environment through inert purge or vacuum,<sup>115</sup> and/or (ii) inducing oxygen vacancy instead of a phase change. In this case, the chemical looping strategy, which allows “tunable” thermodynamic parameters and operating conditions for sub-reactions, enables an equilibrium-limited chemical reaction, which would otherwise be impossible under reasonable operating temperatures.<sup>116</sup> Using a similar principle, other equilibrium-limited reactions such as CO<sub>2</sub> splitting<sup>117</sup> and thermochemical conversions can also be facilitated by CLBC.<sup>93,98,100,118–121</sup>

*d. Tunable exo-/endothermicity of sub-reactions for improved energy integration.* Chemical looping would not alter the enthalpy or entropy changes of the overall reaction under (near) isothermal conditions. However, the distributions of the reaction enthalpy/entropy can be modified by varying the type of the oxygen carrier/redox catalyst and the chemical looping sub-reactions. As such, the heat release/demand of individual chemical looping reactors can be adjusted. This additional degree of freedom can enable significantly improved energy integration and hence reduced exergy/energy losses.<sup>122</sup> With an increased understanding of  $P_{\text{O}_2}$ , phase transition, and catalytic properties for the mixed oxide-based oxygen carriers, the tunabilities of oxygen carriers and hence the corresponding sub-reactions are becoming an increasingly attractive route for efficiency improvements. Fig. 5 summarizes the potential advantages of CLBC over conventional approaches.

### 2.3 Reasons for CLBC – the practical drivers

Besides the abovementioned thermodynamic “drivers”, a strong case can be made for CLBC from a practical standpoint.

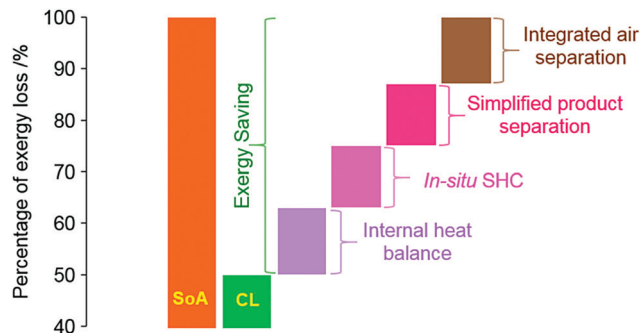


Fig. 5 Schematic illustration of the exergy loss for conventional and chemical looping approaches. SoA: state-of-the-Art (SoA) approaches are typically based on thermal cracking or partial oxidation with oxygen co-feed. SHC refers to selective combustion of hydrogen byproduct, e.g. in CL-ODH, chemical looping oxidative coupling of methane (CL-OCM), and chemical looping dehydroaromatization (CL-DHA).

Commercial implementation of the CLC technology needs to address both technical challenges, in terms of scale-up, and economic challenges, resulting from the low margin of the utility industry and the relatively low economic incentive of the current carbon tax structure.<sup>22,36</sup> A CLC power plant will be more complex and costlier than a standard pulverized coal combustion plant, due to large scale circulating fluidized bed operations, high solids circulation rates, and the need to continuously replenish degraded oxygen carrier particles.<sup>4</sup> Meanwhile, low-cost and long-lifetime oxygen carrier material is required to make CLC cost effective.<sup>4</sup> In addition, concerns over carbon sequestration, a critical downstream step for any carbon capture technology, also limits the wide spread utilization of CLC as a promising carbon capture process.<sup>123–127</sup> CLBC, on the other hand, has the potential to address all the limitations identified above. In contrast to CLC whose economics is likely to be policy driven, CLBC for chemicals production is market driven because the products are of higher value than electric power. As illustrated in Fig. 6, the CLBC concept can be applied to produce chemical products that are significantly more valuable than electricity, thereby increasing the profit margin. In addition, CLBC based on partial oxidation is intrinsically more

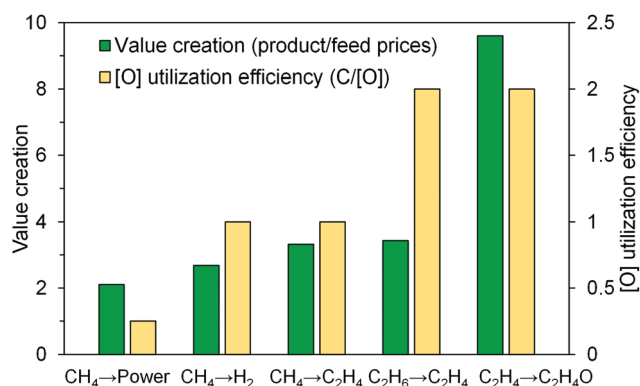


Fig. 6 Valorization and efficiency of oxygen utilization in some typical chemical looping processes.



efficient from an oxygen carrier utilization standpoint. Complete combustion of carbonaceous fuels such as methane consumes a significant amount of lattice oxygen for power generation (four [O] are required for each carbon combusted in CH<sub>4</sub>). In comparison, the same amount of lattice oxygen is able to transform eight carbon atoms in ethane into more valuable ethylene products under a CL-ODH scheme (assuming 100% ethylene selectivity whereas ~90% olefin selectivity has been reported experimentally).<sup>79,81,112,128</sup> If oxygen carriers/redox catalysts with identical oxygen storage capacities are used, the CL-ODH plant would correspond to an eight-fold decrease in solids circulation rate compared to a CLC plant with identical feedstock processing capacity on a weight basis. In addition, at least an order of magnitude increase in reaction kinetics was observed for ethane CL-ODH when compared to methane CLC based on the experience at North Carolina State University.<sup>80,81</sup> Considering that ethylene production could significantly increase the gross margin compared to power generation, the “breakeven” plant size for CL-ODH would likely to be one or more order of magnitude smaller than that for CLC from process economics, reaction stoichiometry, and kinetics standpoints. An even stronger case can be made for CLBC of heavier feedstock/products: applying the chemical looping strategy for butane to butadiene, if successful, would lead to a 16 times higher lattice oxygen utilization efficiency than for methane CLC. This simple case study clearly demonstrates the significant advantages of CLBC from process scale up and economic standpoints. Another interesting point to make is that many CLBC processes themselves do not offer the primary benefit of CLC, *i.e.* CO<sub>2</sub> capture, since the target products are chemicals instead of heat (Table 1). However, the process intensification and second law advantages offered by CLBC can still lead to significant CO<sub>2</sub> reductions by producing less CO<sub>2</sub> per unit amount of product generated. A recent study published by Haribal *et al.*<sup>113</sup> indicated that CL-ODH can reduce the CO<sub>2</sub> emissions for ethylene production by 84% when compared to the commercial ethane cracking processes. Such integrated carbon savings without CO<sub>2</sub> capture (except for the removal of a small amount of CO<sub>2</sub> byproducts *via* conventional stripping) eliminates the needs for geological CO<sub>2</sub> sequestration, a potential technical challenge for any CO<sub>2</sub> capture process including CLC.

### 3. General design strategies for CLBC

#### 3.1 Design considerations for CLBC schemes and limitations

While chemical looping offers various potential advantages as stated in Section 2, it is subjected to a number of limitations, which occasionally are overlooked in CL process analyses. This is somewhat “understandable” considering the complexity of CL processes, which involve dynamic and intertwined gas–solid reactions, *e.g.* the extent of redox catalyst conversion in the reduction step(s) often affects the equilibrium conversion and practical performance of the oxidation step(s). As such, we think it is important to summarize the uniqueness of CLBC as well as its limitations and potential pitfalls in design and

analysis of CLBC schemes. Compared to conventional catalytic oxidation or reduction processes, the chemical looping strategy is capable of:

- (i) Providing inherently separated product streams to eliminate or alleviate the needs for separation;
- (ii) Enhancing process safety by avoiding the direct mixing between a gaseous oxidant (*e.g.* O<sub>2</sub>) and reductant (*e.g.* a hydrocarbon);
- (iii) Redistributing the overall heat of reaction into two or more steps to simplify heat integration;
- (iv) Breaking the “equilibrium barrier” of an overall reaction by carefully selecting redox catalysts, redox pairs<sup>91</sup> and carrying out individual chemical looping steps at substantially different conditions and/or gas–solids contacting modes. For example, the chemical equilibrium limitations in the WGS reaction under a CO–H<sub>2</sub>O co-feed mode can be overcome using the chemical looping strategy over a non-stoichiometric mixed oxide, since such an oxide exhibits a large range of equilibrium  $P_{O_2}$  as a function of oxygen non-stoichiometry.<sup>91</sup> Metcalfe *et al.*<sup>91</sup> investigated such a scheme using a perovskite (La<sub>0.6</sub>Sr<sub>0.4</sub>FeO<sub>3- $\delta$</sub> ) as the oxygen carrier. Using a packed bed reactor operated under a countercurrent contacting mode, a significantly higher overall CO to H<sub>2</sub> conversion (compared to WGS equilibrium conversion) was demonstrated. The unique reactor design and oxygen carrier properties enabled the production of near pure H<sub>2</sub> and CO<sub>2</sub> as separate product streams.
- (v) Potentially achieving higher product selectivity by using lattice oxygen as opposed to gaseous oxidants.

However, one must be aware of the following limitations for chemical looping when designing chemical looping schemes or performing process analyses:

- (i) The sum of Gibbs free energies of the products from all the chemical looping sub-steps should be smaller than that of the feedstock;
- (ii) For a two-step CL process with a single metal/metal oxide redox pair, products from an (oxide) reduction reactor 1 should not be able to oxidize products from an oxidation reactor 2. This principle would apply for both co-currently and counter-currently operated fluidized bed, fixed bed, and moving bed reactors. Using WSG/RWGS reactions<sup>129</sup> as an example, one cannot produce a substantially pure stream of CO<sub>2</sub> or H<sub>2</sub>O from reactor 1 while also producing a substantially pure stream of H<sub>2</sub> or CO from reactor 2 if a single redox pair is used. We have to make a choice between a lower CO<sub>2</sub>/H<sub>2</sub>O yield in reactor 1 and a decreased CO/H<sub>2</sub> yield from reactor 2. Note that this is different from the case illustrated by Metcalfe *et al.*, in which a non-stoichiometric metal oxide was used as the carrier;<sup>91</sup>

(iii) A three-step CL process can allow products from reactor 1 to oxidize products from reactor 2 (since the two reactors and redox pairs do not form a complete loop). However, this is only feasible with an oxidant in step 3 that, when combined with the products from step 2, exhibits a higher equilibrium oxygen partial pressure ( $P_{O_2}$ ) than do the products from step 1;

(iv) Kinetically, chemical looping reactions are generally much slower than conventional catalytic oxidation/reduction reactions. This is due to the fact that the rates of redox reactions





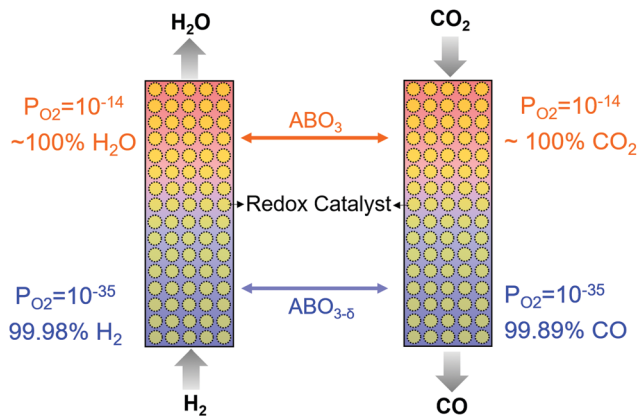


Fig. 7 RWGS in a counter-currently operated two-step chemical looping scheme using  $\text{ABO}_3$  perovskite as a redox catalyst at  $500^\circ\text{C}$ .

are limited by the removal and replenishment of lattice oxygen from/to the bulk crystal structure of the redox catalyst. One exception is the CL-ODH operated at thermal cracking temperatures ( $> 700^\circ\text{C}$ ). In that case, the rate of hydrogen combustion by the redox catalysts can often match or exceed the hydrogen generation rate from the cracking reactions.<sup>80,81</sup>

We note that many mixed oxide-based redox catalysts, *e.g.* perovskites, can accommodate significant lattice defects prior to phase transition.<sup>129,130</sup> As such, the redox thermodynamics of these redox catalysts are unique and cannot be categorized as redox pairs. That is, the equilibrium oxygen partial pressure of the oxide is a continuous function of the oxide's (oxygen) vacancy concentration without incurring a distinct phase change.<sup>131</sup> The unique redox properties of these mixed oxides, coupled with counter-current and/or multiple step chemical looping reactor design, can lead to interesting process configurations that are not subjected to limitations (ii) and (iii) presented above. Fig. 7 illustrates a hypothetical case where a mixed oxide redox catalyst with varying equilibrium  $P_{\text{O}_2}$ s as a function of oxygen storage capacity can be used to break the RWGS equilibrium in a counter-currently operated two-step chemical looping scheme. A comparable effect can be achieved by using more than one redox pairs with different equilibrium  $P_{\text{O}_2}$ s. In reality, however, one must consider the complex relationships between the oxygen non-stoichiometry of the mixed oxide, which dictates the oxygen mass balance of the redox reactions, and the equilibrium  $P_{\text{O}_2}$ , which determines the redox thermodynamics, in addition to kinetic and transport effects.

### 3.2 Oxygen carrier design

**3.2.1 Thermodynamic prerequisites and second law guided oxygen carrier selection.** Although surface properties of the redox catalysts are essential for their reactivity and product selectivity in CLBC, the redox catalysts need to satisfy certain thermodynamic prerequisites. In terms of partial oxidation reactions, the oxide-based redox catalysts need to provide adequate oxygen chemical potential or equilibrium  $P_{\text{O}_2}$  to facilitate product generation. Considering the large material design space for redox catalysts, this thermodynamic prerequisite can be used as an effective tool to narrow down the candidate materials. As

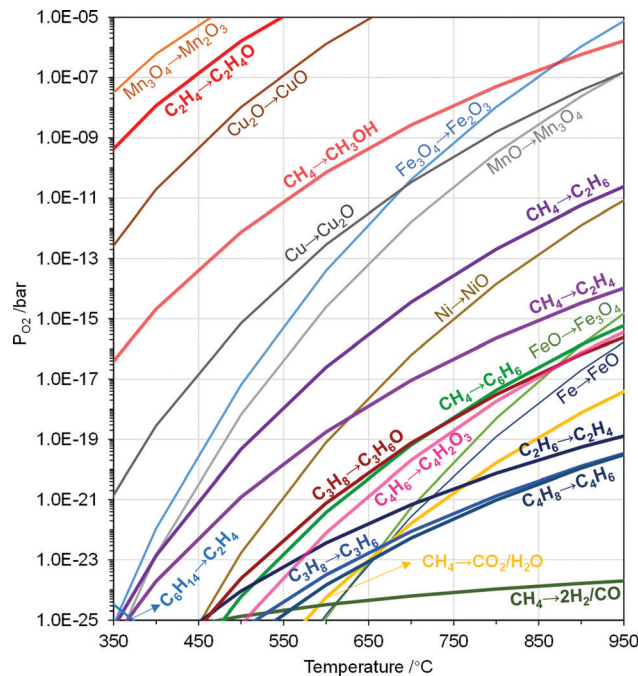


Fig. 8 Modified Ellingham diagram of metal oxide redox and POx reactions.  $\text{MeO}_x \rightarrow \text{MeO}_y$  and hydrocarbons POx reactions correspond to the  $P_{\text{O}_2}$  when  $\Delta G = 0$ . Reactions in pathways of  $\text{MeO}_x \rightarrow \text{MeO}_y$  and hydrocarbons POx are given in Table 2.

illustrated in the Ellingham diagram in Fig. 8, one should consider redox pairs above the equilibrium lines of the corresponding POx (selective oxidation) reaction. Taking oxidative coupling of methane (OCM) as an example, it is apparent that redox pairs such as  $\text{FeO}/\text{Fe}$  and  $\text{Fe}_3\text{O}_4/\text{FeO}$  should not be considered because they would not be sufficiently "oxidative" to facilitate chemical looping OCM at temperatures ranging from  $650^\circ\text{C}$  to  $950^\circ\text{C}$ . On the other hand, redox catalysts composed of  $\text{MnO}_2/\text{Mn}_3\text{O}_4$  or  $\text{Mn}_3\text{O}_4/\text{Mn}_2\text{O}_3$  redox pairs would likely need surface modifications to ensure high selectivities, since their  $P_{\text{O}_2}$ s are well above that required for OCM. Another thermodynamic consideration for redox pair selection is the upper limit for  $P_{\text{O}_2}$ . For a CLBC system operated at near atmospheric pressures, practical requirement of  $> 90\%$   $\text{O}_2$  conversion in the air regeneration step would impose an upper limit of 0.02 bar for equilibrium  $P_{\text{O}_2}$ .

The thermodynamic criteria discussed above are intended for narrowing down the design space of the redox catalysts. Possessing suitable thermodynamic properties alone typically would not ensure high product selectivity. A rare exception is synthesis gas generation. As illustrated in Fig. 2b, synthesis gas is one of the thermodynamically favored oxidation product except for  $\text{CO}_2$  and water. As such, synthesis gas selectivity and yield can be optimized by tuning the  $P_{\text{O}_2}$ s of redox catalysts using Gibbs free energy minimization, as illustrated in a few recent publications.<sup>49,66,67,132</sup> For other POx products with higher Gibbs free energies, however, such a thermodynamically based strategy would not work since the spontaneous decomposition or oxidation of oxygenates to synthesis gas and/or  $\text{CO}_2/\text{H}_2\text{O}$  is both kinetically and thermodynamically favored at elevated temperatures.





Aside from being a redox catalyst selection tool, equilibrium  $P_{O_2}$ s illustrated in the Ellingham diagram can be used to manage heat distributions among the redox steps involved in CLBC. Due to the endothermicity in the oxygen release step of redox catalysts, a large fraction of the fuel oxidation reactions with redox catalysts in CLBC are endothermic, even though oxidation of a fuel with oxygen would be exothermic. Oxides with higher  $P_{O_2}$ s tend to decrease the endothermicity in the fuel oxidation step while decreasing the exothermicity of the redox catalyst re-oxidation step. As such, rationally selected redox pairs can facilitate optimal distribution of the reaction heat in the redox steps to simplify heat integrations. In CLC, a bi-metallic Fe–Cu oxygen carrier composed of CuO and Fe<sub>2</sub>O<sub>3</sub> was proposed to shift the endothermic reduction to exothermic or heat neutral.<sup>133,134</sup> This  $P_{O_2}$  tuning strategy has also been applied to CLBC, *e.g.* in ethane CL-ODH.<sup>82,83,128</sup>

Redox thermodynamic data for monometallic oxides are readily available, as illustrated in Fig. 8. However, the number of suitable redox pairs are limited when considering the aforementioned thermodynamic criteria. Mixed oxides such as hexaaluminates, spinels, or perovskites, offer significantly increased tunability in terms of redox properties. Moreover, equilibrium  $P_{O_2}$ s of many mixed oxides are often variable without incurring phase changes, as discussed in Section 3.1. As such, their multidimensional redox behavior can not be illustrated by a simple redox reaction or a curve in the Ellingham diagram. These unique properties make mixed oxides an intriguing option for CLBC, as illustrated by a number of recent publications.<sup>61,70,135</sup> Among the various mixed oxides, perovskite-type oxides represent a large family of materials that is particularly promising for CLBC due to their highly tunable structural, compositional, and redox properties.<sup>136–138</sup> A recent perspective provided a comprehensive discussion on perovskites as redox catalysts for CLBC.<sup>49</sup>

**3.2.2 Surface modifications of redox catalysts.** Similar to selective oxidation reactions facilitated by heterogeneous

catalysts, a key challenge to redox catalysts in CLBC is to avoid the over-oxidation of desirable products. Recent studies have indicated that surface modifications can be an effective strategy to enhance redox catalysts' activity and/or selectivity.<sup>79–82,128,139</sup> Generally speaking, electrophilic (surface) oxygen species such as O<sup>−</sup>, O<sub>2</sub><sup>2−</sup> (peroxide), and O<sub>2</sub><sup>−</sup> (superoxide) are often attributed to non-selective oxidation reactions due to their high activity and electron affinity.<sup>49,140</sup> In comparison, lattice oxygen is often believed to be the “selective” oxygen species in both CLBC and catalytic oxidation reactions involving oxide catalysts. However, such an assignment can be a rather simplistic view since the presence and relative abundance of these oxygen species is not only dependent on the type of oxide but also the reaction conditions and the degree of reduction of the oxide. Recent experimental evidences indicate that lattice oxygen species can be dynamically converted to electrophilic oxygen under chemical looping reaction conditions.<sup>58,139</sup> As such, the abundance of electrophilic oxygen species during the partial oxidation step can be affected by a dynamic balance between: (i) the flux of lattice oxygen from the bulk oxide lattice to the surface; and (ii) the rate of oxygen removal from the surface of the redox catalyst. More facile lattice oxygen flux to the surface would hence allow the transition or relaxation of O<sup>2−</sup> to electrophilic oxygen especially when such oxygen is bonded with high valency transition metal cations, *e.g.* Fe<sup>4+</sup> or Mn<sup>4+</sup>. Therefore, one would anticipate a low product selectivity resulting from the dynamic formation of the electrophilic oxygen species on the surface.

On the other hand, decreased CO<sub>x</sub> formation can be achieved by facilitating more facile surface oxygen removal compared to the lattice oxygen flux since the formation of electrophilic oxygen species can be minimized. To ensure a relatively faster surface oxygen removal rate, one can either decelerate the O<sup>2−</sup> flux or enhance the surface activity of the redox catalyst, as illustrated in Fig. 9. Recent studies have confirmed the effectiveness of both



Fig. 9 Schematic illustration of surface modifications of redox catalysts to improve product selectivity by changing the relative rates of surface oxygen removal and bulk O<sup>2−</sup> flux. The arrows in the three cases (a–c) correspond to the O<sup>2−</sup> fluxes and the colors correspond to the relative intensity of the O<sup>2−</sup> fluxes.



approaches. Gao *et al.*<sup>79,128</sup> investigated a  $\text{La}_x\text{Sr}_{2-x}\text{FeO}_{4-\delta}$ -based redox catalyst for CL-ODH of ethane to ethylene. While  $\text{La}_x\text{Sr}_{2-x}\text{FeO}_{4-\delta}$  promoted the complete combustion of ethane, alkali metal oxide promoters enriched on the surface of the mixed oxide and significantly inhibited the  $\text{O}^{2-}$  flux from the bulk. As a result, 90% ethylene selectivity was achieved. Besides alkali metal oxide promoters, alkali salt modification also play a similar role to depress  $\text{O}^{2-}$  flux of the redox catalyst for the selective combustion of hydrogen (SHC) in CL-ODH of ethane to ethylene.<sup>81–83</sup> Enhancement of redox catalyst's surface activity was reported for methane partial oxidation for synthesis gas generation. Shafiefarhood *et al.*<sup>139</sup> reported that impregnating Rh on a  $\text{CaMnO}_3$ -based redox catalyst decreased the methane activation energy on the oxide surface by more than 95%, leading to a decrease in methane conversion temperature by 300 °C and a significantly increased synthesis gas selectivity. Those practices reveal that tuning of the oxygen transfer rate from bulk of a redox catalyst to the catalytically active species on the surface is of great importance for POx. In a redox-involved context, Machida at Kumamoto University found that surface  $\text{CeO}_2$  could act as a gateway for oxygen storage in and release from various oxides acting as an oxygen reservoir, such as  $\text{Fe}_2\text{O}_3$  or  $\text{La}_2\text{O}_2\text{SO}_4$ .<sup>141–143</sup>

Although the abovementioned strategy for catalyst selectivity optimization has been shown to be effective, we note that it still represents a rather simplified view of the redox catalyst. It would be desirable, from both scientific and engineering standpoints, to determine the active sites and elementary reaction pathways of the chemical looping reactions, but such tasks can be daunting due to the highly dynamic nature of chemical looping reactions. In fact, both the bulk and surface of the redox catalysts can dynamically rearrange throughout the redox cycle.<sup>144</sup> The utilization of modern catalysis and surface science techniques in a few recent studies have already started to reveal fundamental insights on the mechanisms of the redox reactions.<sup>78,128,145–151</sup> However, significant progress still needs to be made to gain additional fundamental insights and this should be a topic of focus for redox catalyst studies within the foreseeable future.

In principle, one can also combine heterogeneous catalysts or active sites with an oxygen storage material to achieve combined catalytic and chemical looping functions. For instance, as will be discussed in Section 4.2.2, Chan *et al.* demonstrated the CL epoxidation of ethylene over  $\text{Ag}/\text{SrFeO}_3$  redox catalyst where Ag acts as the epoxidation catalyst and  $\text{SrFeO}_3$  acts as the oxygen storage material for oxygen donation.<sup>78</sup> Ethylene was converted to ethylene oxide (with  $\text{CO}_2$  and  $\text{H}_2\text{O}$  by products) in the reduction step and the reduced redox catalyst was then replenished with air. Many other CLBC systems can potentially take advantage of such a strategy. Examples include alkane ODH, methane dehydroaromatization, methane oxidative coupling, among others. These topics are discussed in Sections 4.1 and 4.2.

We note that the above discussions are intended to cover the key strategies and the underlying principles for the design of CLBC redox catalysts and reaction schemes. From a practical standpoint, the redox catalyst's activity, available oxygen storage capacity, stability, mechanical strength,<sup>152,153</sup> heat management,<sup>154</sup>

cost and environmental risk should all be taken into account to optimize the redox catalyst. The higher margin in the chemical industry over the utility industry can afford significantly higher redox catalyst cost in CLBC. While the raw material cost of some redox catalysts can be high (*e.g.* ceria-based materials), the additional value created for chemical production can potentially accommodate higher redox catalyst costs. Under CLBC schemes, redox catalysts functions similar to heterogeneous catalysts in chemical production processes.<sup>155</sup> It is further noted that a number of CLBC redox catalysts have shown chemical productivities (in terms of weight hourly space velocity) comparable to commercial catalysts.<sup>111</sup> The aforementioned factors should also be considered along with reactor design parameters, operating conditions as well as the reactor/process economics for CLBC process development.

## 4. CLPOx/CLBC process schemes

### 4.1 Chemical looping for $\text{C}_1$ valorization

$\text{CH}_4$  is the main constituent of natural gas and an important feedstock for the production of commodity chemicals and liquid fuels.<sup>156</sup> It can be converted into methanol,  $\text{NH}_3$  and other higher hydrocarbons either directly *via* a one-step process or indirectly *via* a synthesis gas (a mixture of  $\text{CO}$  and  $\text{H}_2$ ) intermediate. At present, the only cost-effective route to convert  $\text{CH}_4$  into valuable chemicals is *via* synthesis gas.<sup>157</sup> Synthesis gas can be obtained from  $\text{CH}_4$  using the following three reactions: steam reforming (R62), dry reforming (R63) and partial oxidation (R49).<sup>158</sup> At the industrial scale, the catalytic steam methane reforming (SMR) is used predominantly to produce synthesis gas. Although SMR is highly optimized, it is a very energy-intensive process and releases large quantities of  $\text{CO}_2$  into the atmosphere.<sup>131</sup> To reduce the energy demand of SMR, a part of  $\text{CH}_4$  is combusted internally with oxygen or air. The resulting so-called autothermal reforming process enhances  $\text{CH}_4$  conversion and reduces  $\text{CO}_2$  emissions, but at the expense of lowered synthesis gas yields.<sup>57</sup> Recently, dry reforming of methane (DRM) has received a lot of attention due to its potential to convert two greenhouse gases (*i.e.*  $\text{CH}_4$  and  $\text{CO}_2$ ) into an equimolar mixture of  $\text{CO}$  and  $\text{H}_2$  desired for Fischer–Tropsch synthesis.<sup>159,160</sup> Although this concept has many environmental and economic incentives, there is currently no commercial process for DRM. Unlike reforming reactions, methane POx is mildly exothermic allowing for auto-thermal operation at temperatures exceeding 1000 °C. Methane POx can be performed either at elevated temperatures (> 1200 °C) without a catalyst or at lower temperatures (< 900 °C) in the presence of a platinum group metal (PGM) based catalyst (*e.g.*, Pt, Pd, Rh, Ru, *etc.*).<sup>161,162</sup> Commercial methane POx processes incur a significant efficiency loss due to high operating temperatures. The catalytic route can be costly due to the high cost of the PGM and safety concerns. Both approaches also are challenged by the high cost and energy intensity of pure  $\text{O}_2$ .

In 2000, Lyon and Cole<sup>163</sup> proposed a chemical looping approach to transfer oxygen from air to a fuel using solid oxygen carriers



Table 2 Reactions for  $\text{MeO}_x \rightarrow \text{MeO}_y$ , hydrocarbon partial oxidation, and calcium/carbonate looping reactions illustrated in Fig. 8

Category	Pathway	Reaction
Metal oxide redox pairs	$\text{Mn}_3\text{O}_4 \rightarrow \text{Mn}_2\text{O}_3$	(R42) $4\text{Mn}_3\text{O}_4 + \text{O}_2(\text{g}) \rightarrow 6\text{Mn}_2\text{O}_3$
	$\text{MnO} \rightarrow \text{Mn}_3\text{O}_4$	(R43) $6\text{MnO} + \text{O}_2(\text{g}) \rightarrow 2\text{Mn}_3\text{O}_4$
	$\text{FeO} \rightarrow \text{Fe}_3\text{O}_4$	(R44) $6\text{FeO} + \text{O}_2(\text{g}) \rightarrow 2\text{Fe}_3\text{O}_4$
	$\text{Cu}_2\text{O} \rightarrow \text{CuO}$	(R45) $2\text{Cu}_2\text{O} + \text{O}_2(\text{g}) \rightarrow 4\text{CuO}$
	$\text{Cu} \rightarrow \text{CuO}$	(R46) $2\text{Cu} + \text{O}_2(\text{g}) \rightarrow 2\text{CuO}$
	$\text{Ni} \rightarrow \text{NiO}$	(R47) $2\text{Ni} + \text{O}_2(\text{g}) \rightarrow 2\text{NiO}$
	Combustion	$\text{CH}_4 \rightarrow \text{Power}$
Partial oxidation/selective oxidation		(R49) $2\text{CH}_4(\text{g}) + \text{O}_2(\text{g}) \rightarrow 2\text{CO}(\text{g}) + 4\text{H}_2(\text{g})$ (R50) $2\text{CH}_4(\text{g}) + \text{O}_2(\text{g}) \rightarrow 2\text{CH}_3\text{OH}(\text{g})$ (R51) $2\text{C}_2\text{H}_4(\text{g}) + \text{O}_2(\text{g}) \rightarrow 2\text{C}_2\text{H}_4\text{O}(\text{g})$ (R52) $\text{C}_3\text{H}_8(\text{g}) + \text{O}_2(\text{g}) \rightarrow \text{C}_3\text{H}_6\text{O}(\text{g}) + \text{H}_2\text{O}(\text{g})$ (R53) $2/5\text{C}_4\text{H}_6(\text{g}) + \text{O}_2(\text{g}) \rightarrow 2/5\text{C}_4\text{H}_2\text{O}_3(\text{g}) + 4/5\text{H}_2\text{O}(\text{g})$
Oxidative coupling	$\text{CH}_4 \rightarrow \text{C}_2\text{H}_4$	(R54) $2\text{CH}_4(\text{g}) + \text{O}_2(\text{g}) \rightarrow \text{C}_2\text{H}_4(\text{g}) + 2\text{H}_2\text{O}(\text{g})$ (R55) $4\text{CH}_4(\text{g}) + \text{O}_2(\text{g}) \rightarrow 2\text{C}_2\text{H}_6(\text{g}) + 2\text{H}_2\text{O}(\text{g})$ (R56) $4/3\text{CH}_4(\text{g}) + \text{O}_2(\text{g}) \rightarrow 2/9\text{C}_6\text{H}_6(\text{g}) + 2\text{H}_2\text{O}(\text{g})$
	Oxidative dehydroaromatization (DHA) Oxidative dehydrogenation or cracking	(R57) $2\text{C}_2\text{H}_6(\text{g}) + \text{O}_2(\text{g}) \rightarrow 2\text{C}_2\text{H}_4(\text{g}) + 2\text{H}_2\text{O}(\text{g})$ (R58) $2\text{C}_3\text{H}_8(\text{g}) + \text{O}_2(\text{g}) \rightarrow 2\text{C}_3\text{H}_6(\text{g}) + 2\text{H}_2\text{O}(\text{g})$ (R59) $2\text{C}_4\text{H}_8(\text{g}) + \text{O}_2(\text{g}) \rightarrow 2\text{C}_4\text{H}_6(\text{g}) + 2\text{H}_2\text{O}(\text{g})$ (R60) $2\text{C}_6\text{H}_{14}(\text{g}) + \text{O}_2(\text{g}) \rightarrow 6\text{C}_2\text{H}_4(\text{g}) + 2\text{H}_2\text{O}(\text{g})$ (R61) $2\text{C}_6\text{H}_{14}(\text{g}) + \text{O}_2(\text{g}) \rightarrow 4\text{C}_3\text{H}_6(\text{g}) + 2\text{H}_2\text{O}(\text{g})$
Reforming	$\text{CH}_4 \rightarrow \text{Synthesis gas}$	(R62) $\text{CH}_4(\text{g}) + \text{H}_2\text{O}(\text{g}) \rightarrow 3\text{H}_2(\text{g}) + \text{CO}(\text{g})$ (R63) $\text{CH}_4(\text{g}) + \text{CO}_2(\text{g}) \rightarrow 2\text{H}_2(\text{g}) + 2\text{CO}(\text{g})$ (R64) $\text{CH}_4(\text{g}) \rightarrow \text{C}(\text{s}) + 2\text{H}_2(\text{g})$ (R65) $6\text{CH}_4(\text{g}) \rightarrow \text{C}_6\text{H}_6(\text{g}) + 9\text{H}_2(\text{g})$ (R66) $\text{CH}_4(\text{g}) + \text{H}_2\text{O}(\text{g}) \rightarrow \text{C}_2\text{H}_5\text{OH}(\text{g}) + \text{H}_2(\text{g})$ (R67) $\text{CH}_4(\text{g}) + 3\text{CO}_2(\text{g}) \rightarrow 4\text{CO}(\text{g}) + 2\text{H}_2\text{O}(\text{g})$ (R68) $\text{CaO}(\text{s}) + \text{CO}_2(\text{g}) \leftrightarrow \text{CaCO}_3(\text{s})$ (R69) $\text{CH}_4(\text{g}) + 2\text{H}_2\text{O}(\text{g}) + \text{CaO} \rightarrow \text{CaCO}_3(\text{s}) + 4\text{H}_2(\text{g})$ (R70) $2\text{CH}_4(\text{s}) + 4\text{CuO}(\text{s}) + 4\text{NiO}(\text{s}) + \text{CaCO}_3(\text{s}) \rightarrow \text{CaO}(\text{s}) + 4\text{Cu}(\text{s}) + 4\text{Ni}(\text{s}) + 3\text{CO}_2(\text{g}) + 4\text{H}_2\text{O}(\text{g})$
RWGS	$\text{CO}_2/\text{H}_2 \rightarrow \text{CO}/\text{H}_2\text{O}$	(R71) $\text{CO}_2(\text{g}) + \text{H}_2(\text{g}) \rightarrow \text{CO}(\text{g}) + \text{H}_2\text{O}(\text{g})$
	Methanol oxidation	(R72) $2\text{CH}_3\text{OH}(\text{g}) + \text{O}_2(\text{g}) \rightarrow 2\text{CH}_2\text{O}(\text{g}) + 2\text{H}_2\text{O}(\text{g})$

for a number of applications such as pollution control, production of inert gases, delivery of heat to endothermic reactions, *etc.* Since then various chemical looping cycles have been proposed to address the limitations of conventional  $\text{CH}_4$  conversion processes (see in Table 2). Sections 4.1.1 and 4.1.2 provide a brief summary of the chemical looping approaches related to indirect and direct  $\text{CH}_4$  conversion routes, respectively.

**4.1.1 CL methane conversion to synthesis gas.** Compared to conventional  $\text{CH}_4$  reforming processes that utilize gaseous oxidants, the two-step chemical looping approach, schematized in Fig. 10, relies on the lattice oxygen of a solid oxygen carrier to partially oxidize  $\text{CH}_4$  *via* (R3).<sup>63,164,165</sup> A subsequent re-oxidation step then replenishes the lattice oxygen. If air is used in the regeneration step to re-oxidize the reduced oxygen carrier, *via* (R4), the overall reaction is methane POx and hence the chemical looping cycle is also referred to as CL-methane POx.<sup>166</sup> Unlike the conventional methane POx reaction (R49), the chemical looping route does not require pure oxygen, thereby eliminating the need of an energy- and cost-intensive air separation unit (ASU). Further, the actual  $\text{CH}_4$  conversion step using lattice oxygen (when the oxygen carrier is reduced) is endothermic. Depending upon the thermodynamic characteristics of the oxygen carrier, it can also be re-oxidized using  $\text{H}_2\text{O}$ ,<sup>167,168</sup> see (R5), resulting in the so-called CL-SMR.<sup>64,169</sup> Here, two separate streams of synthesis gas (with a  $\text{CO}:\text{H}_2$  ratio of 1:2) and high-purity  $\text{H}_2$  are produced inherently, thus intensifying the

production of high-purity  $\text{H}_2$  from  $\text{CH}_4$ . It is noted here that CL-SMR would become the CL-WGS process if  $\text{CO}$  instead of  $\text{CH}_4$  were used to reduce the oxygen carrier.<sup>42,96,170,171</sup> Both CL-SMR and CL-WGS are also referred to as CL-water-splitting processes.<sup>47,172</sup> Recently the CL-DRM cycle, in which  $\text{CO}_2$  is used to regenerate the oxygen carrier (*via* (R6)), has been proposed to maximize the conversion of  $\text{CO}_2$ , captured from various emission sources, to  $\text{CO}$ .<sup>96,162,167,173–175</sup> The modified CL-methane POx coupled with  $\text{H}_2\text{O}-\text{CO}_2$  splitting<sup>176</sup> for the oxygen carrier regeneration allows the production of syngas with identical compositions in both the reduction and regeneration steps.<sup>121,177,178</sup>

The oxygen carriers used in the above-mentioned chemical looping processes are often called redox catalysts because they simultaneously act as an oxygen donor (*i.e.* as a reactant) and as a catalyst.  $\text{NiO}$ ,  $\text{Fe}_2\text{O}_3$ ,  $\text{CeO}_2$ , spinel and certain perovskite-based oxides possess suitable thermodynamic properties that permit their use as redox catalysts for chemical looping conversion of  $\text{CH}_4$  to synthesis gas.<sup>41,179–184</sup> Among various oxygen carriers, Ni-based oxygen carriers are particularly attractive for CLR owing to their high catalytic activity for  $\text{CH}_4$  reforming.<sup>56,185–191</sup>  $\text{NiO}$  possesses a low selectivity towards synthesis gas, but as  $\text{NiO}$  reduces to  $\text{Ni}$  the selectivity towards synthesis gas increases.<sup>56</sup> However,  $\text{Ni}$  is highly prone to deactivation *via* carbon deposition under low oxygen partial pressures<sup>187</sup> and the formation of sulfides in the presence of feedstock containing sulfur species.<sup>192</sup>





Fig. 10 Schematic and reaction schemes of different CL cycles for conversion of  $\text{CH}_4$  to synthesis gas.

Moreover, Ni is a known carcinogenic, thus possessing significant health and safety risks.<sup>193</sup> Fe-Based redox catalysts are also very promising for CL-methane POx owing to their suitable thermodynamic properties for both  $\text{CH}_4$  reforming and  $\text{H}_2\text{O}/\text{CO}_2$  splitting, low cost and minimal environmental impact.<sup>90,194</sup> However, compared to Ni-based oxygen carriers, Fe-based redox catalysts possess a low reactivity with  $\text{CH}_4$  and a low selectivity towards synthesis gas.<sup>195</sup> Because of their high activity and thermal stability, La-based perovskites with Fe, Mn, or Co as B-site cations have been widely investigated in recent years for CL-methane POx and CL-SMR.<sup>58–60,64,66,88,89,139,179,196–198</sup> As mentioned above, the structural and compositional tailoring of the redox catalysts can open up extensive possibilities for improvements of CLR processes. Since the objective of this part of the perspective is not to present a comprehensive review of the redox catalysts used for CLR, we briefly summarize some of the strategies that have been demonstrated to enhance the reactivity and selectivity of redox catalysts in the following sections.

**Stabilization by a mixed conductive support.** The redox catalysts typically comprise of a primary metal oxide as an oxygen reservoir and an inert support to stabilize the metal oxide. The appropriate combinations of oxygen carriers and supports can potentially increase the selectivity of synthesis gas and reduce catalyst deactivation.<sup>199</sup> For example, it has been demonstrated that the use of a mixed conductive support (such as  $\text{La}_{0.8}\text{Sr}_{0.2}\text{FeO}_{3-\delta}$  or  $\text{Ca}_{0.8}\text{Sr}_{0.2}\text{Ti}_{0.8}\text{Ni}_{0.2}\text{O}_3$ ) enables fast solid state  $\text{O}^{2-}$  and electron/hole exchange<sup>92,131,200</sup> between the metal oxide crystallites and the surface of oxygen carrier particle, thereby allowing effective oxygen removal and restoration during reactions (R3) and (R4).<sup>201,202</sup> Importantly, supports that promote the outwards diffusion of active metal should be avoided because such supports can potentially deactivate the redox catalysts *via* surface enrichment and agglomeration of the active metal oxide crystallites.<sup>203</sup>

**Core-shell architecture.** The spatial distribution of support in an oxygen carrier particle also significantly affects its activity

and selectivity.<sup>204</sup> For example, Shafiefarhood *et al.*<sup>59</sup> demonstrated that an  $\text{Fe}_2\text{O}_3@ \text{La}_x\text{Sr}_{1-x}\text{FeO}_3$  (LSF) core-shell material was 10–200 times more active, selective and resistant to carbon formation than composite iron oxide-based materials comprising inert ( $\text{Al}_2\text{O}_3$  and  $\text{MgAl}_2\text{O}_4$ ), ionic-conductive (YSZ), and mixed-conductive (LSF) supports. Neal *et al.*<sup>58,60</sup> and Shafiefarhood *et al.*<sup>205</sup> investigated the mechanism for  $\text{CH}_4$  partial oxidation over an  $\text{Fe}_2\text{O}_3@ \text{LSF}$  core-shell material and found four distinct regimes, *viz.* (i) full oxidation of  $\text{CH}_4$  to  $\text{CO}_2$ , (ii) competing full oxidation to  $\text{CO}_2$  and partial oxidation to CO, (iii) partial oxidation of  $\text{CH}_4$  to CO, and (iv)  $\text{CH}_4$  decomposition. During deep oxidation in region (i) “loose” lattice oxygen from the iron oxide core transports and evolves into surface oxygen species. In region (ii), both selective and non-selective (*i.e.* loose) oxygen species compete for  $\text{CH}_4$  oxidation, with  $\text{O}^{2-}$  flux and overall oxygen availability determining the product ratio. In region (iii), dissociatively adsorbed  $\text{CH}_x$  species are partially oxidized by  $\text{O}^{2-}$  species on the surface, resulting in an increase in metallic iron species along with a decrease in the concentration of oxygen surface species. Coke formation in region (iv) results from the depletion of  $\text{O}^{2-}$  in the iron oxide core and  $\text{CH}_4$  decomposition over metallic iron precipitated from the LSF lattice. Transient pulse studies showed that oxygen anion conduction and/or its evolution to electrophilic surface oxygen species is the rate-limiting step in all the reduction regions of interest.<sup>205</sup> Oxygen atoms maintain a modified Mars-van Krevelen (MvK) mechanism throughout the reaction, while the mechanism of  $\text{CH}_4$  oxidation changes from the Eley-Rideal in region (i) to a Langmuir-Hinshelwood-like mechanism in region (iii). These findings indicate that redox catalyst architectures that inhibit the formation of non-selective surface oxygen species while maintaining a steady supply of lattice oxygen to the surface can potentially lead to an improved performance for CL-methane POx.

**Use of promoters.** Ni- and Fe-based oxygen carriers have a high catalytic activity for the decomposition of  $\text{CH}_4$ , *via* (R64). Promotion of redox catalysts with a transition metal that is inactive for the dissociation of  $\text{CH}_4$ , *e.g.* Cu, Ag, or Au, can suppress the formation of carbon during the  $\text{CH}_4$  reforming step.<sup>206</sup> In this regard, Imtiaz *et al.*<sup>207</sup> demonstrated that doping of  $\text{Fe}_2\text{O}_3$  with CuO significantly lowers its propensity for coke deposition in a  $\text{CH}_4$  atmosphere. The high resistance to carbon deposition of Cu modified oxygen carriers was found to be due to the (partial) coverage of surface Fe with Cu. Some transition metals (*e.g.* Ru, Rh, Ir, *etc.*) can expedite the dissociation of C–H bonds in  $\text{CH}_4$ <sup>206</sup> and can therefore enhance the reforming activity of redox catalysts. Shafiefarhood *et al.*<sup>139</sup> reported a significant improvement in the performance of Rh-promoted  $\text{CaMnO}_3$  and  $\text{La}_2\text{Ce}_2\text{O}_7$  redox catalysts for  $\text{CH}_4$  reforming. It was found that a higher concentration of  $\text{CH}_x$  species on the surface of Rh-promoted catalysts accelerates the extraction of facile oxygen resulting in an enhanced  $\text{O}^{2-}$  conduction through the bulk of the redox catalyst. Therefore, a higher redox activity and synthesis gas selectivity was observed for Rh promoted redox catalysts at temperatures as low as





500 °C than for pristine catalysts. Similar findings were reported by Palchela *et al.*<sup>208</sup> for Rh promoted  $\text{La}_{0.8}\text{Sr}_{0.2}(\text{Fe}_{0.8}\text{Co}_{0.2})_{1-x}\text{Ga}_x\text{O}_3$ , which showed a 40% enhancement in CO selectivity for methane conversion at 600 °C. Noble metals, modification with  $\text{CeO}_2$  nanoparticles can also improve the oxygen storage capacity and mobility in perovskite<sup>68</sup> or iron oxides,<sup>63,64,141,209</sup> which in turn can enhance the availability of lattice oxygen for the methane POx to synthesis gas.

**4.1.2 CL methane to chemicals.** The selective oxidation of  $\text{CH}_4$  to value-added chemicals has been studied extensively, but no significant breakthroughs have been achieved that would enable commercial adoption. A key reason lies in the thermodynamic limitations:  $\text{CH}_4$  is more favorably converted to  $\text{CO}_2$  and  $\text{H}_2\text{O}$  than to  $\text{C}_1$  oxygenates or  $\text{C}_{2+}$  hydrocarbons, as evident when comparing the Gibbs free energy changes for the respective reactions (Fig. 2b).<sup>210</sup> In consequence, the reactions must be controlled kinetically through suitable catalysts.<sup>211</sup> Even if  $\text{CH}_4$  is converted successfully to the desired products, these products tend to be more reactive than  $\text{CH}_4$  and can get oxidized readily to  $\text{CO}_x$ , thus lowering the overall yield.

For CL-based  $\text{CH}_4$  partial oxidation, gaseous  $\text{O}_2$  is substituted by lattice oxygen. This eliminates the expensive air separation, ensures safer operation and inhibits non-selective oxidation reactions in the gas phase when carried out at high temperatures.<sup>212,213</sup> In addition, the use of lattice oxygen is also shown to potentially benefit the product selectivity.<sup>214,215</sup> Challenges in the design of suitable redox catalysts relate to the following aspects of CL-based  $\text{CH}_4$  oxidation processes:<sup>213</sup> (i) CL would likely to require higher operating temperature than co-feed which may negate the selectivity benefit of using lattice oxygen, and (ii) the surface and bulk of the CL redox catalyst tend to undergo notable changes during the redox reactions, which makes maintaining a high product selectivity challenging. Below we briefly discuss several  $\text{CH}_4$  conversion schemes that were shown to be feasible in chemical looping mode.

**Oxidative coupling.** The oxidative coupling of methane (OCM) is a method proposed in the early 1980s to convert  $\text{CH}_4$  directly into ethane and ethylene according to reactions (R55) and (R54) at temperatures  $> 600$  °C.<sup>216</sup> The exothermic reactions can be carried out either in “normal” catalytic fashion where oxygen is co-fed with  $\text{CH}_4$ , or in a cyclic redox (chemical looping) operation expressed by reactions (R7) and (R8).<sup>212,217,218</sup> For either mode, an upper limit of 25–30% yield for  $\text{C}_{2+}$  hydrocarbon products has been supposed based on mechanistic considerations,<sup>215,217,219</sup> which is near the yields required for economic competitiveness.<sup>220–222</sup> Various engineering means have been suggested to improve yields, but no breakthroughs have been achieved to date.<sup>213,223,224</sup> It is worth noting that early studies by Atlantic Richfield company, which operated a 12 year program on OCM focusing on the redox mode, reported an approximately 10% higher  $\text{C}_{2+}$  selectivity under the chemical looping mode compared to the conventional catalytic route.<sup>215</sup>

The OCM reaction occurs *via* a complex mechanism including coupled heterogeneous and homogeneous steps (depending on the catalyst) and is initiated by breaking one of the four

equivalently strong C–H bonds of the methane molecule to form methyl radicals.<sup>224,225</sup> The low yields for  $\text{C}_{2+}$  hydrocarbons obtained so far are largely due to secondary oxidation reactions of these reactive products with oxygen weakly bound to the surface of the catalyst and with oxygen in the gas phase, forming a significant amount of  $\text{CO}_x$ .<sup>71,222,226,227</sup> More recently, the upper yield limit for  $\text{C}_{2+}$  products in the absence of gas phase reactions was estimated to be as high as 60% at elevated pressure, which could possibly be achieved *via* chemical looping.<sup>225,228</sup> Experiments with doped- $\text{Mg}_6\text{MnO}_8$  and  $\text{Mn}/\text{Na}_2\text{WO}_4/\text{SiO}_2$ , a catalyst previously shown to attain single pass  $\text{C}_{2+}$  product yields of ~25% in co-feed mode,<sup>229,230</sup> gave  $\text{C}_{2+}$  product yields of 23.2% and 25%, respectively under a chemical looping mode.<sup>231–234</sup> In these studies, the selectivity toward the coupling reaction was high, but the  $\text{H}_2$  molecules formed through the dehydrogenation reaction of ethane competed with  $\text{CH}_4$  molecules for reacting with activated oxygen on the surface of the catalyst, resulting in a decreasing rate of  $\text{CH}_4$  activation and consequently lower  $\text{C}_{2+}$  yields than expected.<sup>222</sup> Hence, improving the activity of the redox catalysts without negatively affecting their selectivity is one important task. In addition, the reaction conditions (*e.g.* temperature and pressure) probably need to be revised to surpass  $\text{C}_{2+}$  product yields of 30%.<sup>235</sup> The importance of dopants in  $\text{Mg}_6\text{MnO}_8$  was highlighted recently,<sup>234</sup> and it was found that Li-doping-induced oxygen vacancies reduce the adsorption energy of methyl radicals and increase the C–H activation barrier, resulting in an increased selectivity towards  $\text{C}_{2+}$ . For doped  $\text{Na}_2\text{WO}_4$  on  $\text{SiO}_2$ -supported  $\text{MO}_x$  catalysts (where  $\text{M} = \text{V}, \text{Cr}, \text{Mn}, \text{Fe}, \text{Co}$  or  $\text{Zn}$ ) it was found previously that their performance is correlated with their electrical conductivity.<sup>236</sup>

**Non-oxidative dehydroaromatization.** Non-oxidative dehydroaromatization (DHA) is a promising route to produce  $\text{C}_6\text{H}_6$  directly from  $\text{CH}_4$ . Conversion of  $\text{CH}_4$  to  $\text{C}_6\text{H}_6$  *via* reaction (R65) was first reported in 1993 using a ZSM-5 zeolite modified with Mo or Zn.<sup>237</sup> Since then, a wide range of metals and zeolites have been investigated for DHA of  $\text{CH}_4$ , but Mo-modified ZSM-5-based catalysts still remain the most effective.<sup>238,239</sup>

Despite being a potentially attractive pathway to selectively produce high-value aromatics,  $\text{CH}_4$  DHA has not yet been commercialized due to the following reasons: Firstly,  $\text{CH}_4$  DHA (R65) is endothermic and limited by the thermodynamic equilibrium. From Fig. 11, it can be seen that the amount of  $\text{C}_6\text{H}_6$  produced is only significant at high temperatures (*i.e.*  $> 600$  °C). Secondly, the inevitable decomposition of  $\text{CH}_4$  (R64) under reaction conditions not only lowers the equilibrium yield of  $\text{C}_6\text{H}_6$  substantially but also causes coking and catalyst deactivation.<sup>241</sup> Consequently,  $\text{C}_6\text{H}_6$  selectivity is typically between 60–80% at  $\text{CH}_4$  conversions of ~10%, corresponding to net  $\text{C}_6\text{H}_6$  yields of  $< 10\%$  under conditions of practical interest.<sup>239</sup> Several strategies have been proposed to overcome these barriers, *e.g.* the use of membrane reactors to remove  $\text{H}_2$  produced during reaction<sup>242</sup> or the periodic regeneration of the coked catalyst.<sup>73</sup> Recently, a CL approach was suggested to potentially address all three limitations of conventional  $\text{CH}_4$  DHA process as following (Fig. 11):<sup>145</sup> (i) the reactive separation



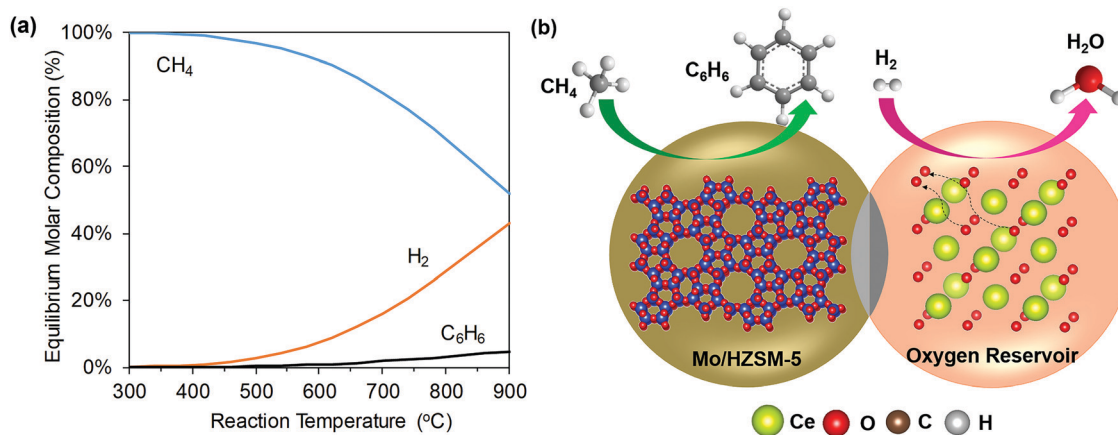


Fig. 11 (a) Equilibrium product composition of CH<sub>4</sub> DHA as a function of reaction temperature. (b) A potential configuration of methane CL-DHA process (the air regeneration step is not shown).<sup>240</sup>

of H<sub>2</sub> from the products of DHA reaction to enhance the equilibrium CH<sub>4</sub> conversion and C<sub>6</sub>H<sub>6</sub> yield, (ii) heat generation by selective H<sub>2</sub> combustion to decrease the overall heat demand, and (iii) *in situ* steam generation to reduce coking. However, care must be taken because excessive presence of steam can negatively affect catalyst selectivity. Moreover, re-oxidation requirements for CL may affect the active MoC<sub>x</sub> species on Mo/ZSM-5. Using Mo/HZSM-5 (as the DHA catalyst) and Ce<sub>0.9</sub>Gd<sub>0.1</sub>O<sub>y</sub> (as the SHC redox catalyst) in a composite bed, an approximately three times higher aromatics yield was demonstrated compared to the conventional process employing only Mo/HZSM-5 without periodic regeneration with oxygen.<sup>145</sup>

Besides the two-step approach, Brady *et al.*<sup>240</sup> proposed a four-step approach for CL-DHA. In the first step, DHA reaction is performed on a Mo/H-ZSM-5 catalyst at 700 °C to obtain a mixture of CH<sub>4</sub>, C<sub>6</sub>H<sub>6</sub> and H<sub>2</sub>. Subsequently, H<sub>2</sub> present in the DHA effluent is combusted selectively using Fe<sub>3</sub>O<sub>4</sub>. The H<sub>2</sub>O produced in step two is then removed using a high-temperature water adsorbent, *e.g.* Zeolite 5A, which can be regenerated using a temperature swing. Finally, the reduced oxygen carrier is re-oxidized back to Fe<sub>3</sub>O<sub>4</sub> using steam. To increase the conversion of CH<sub>4</sub> and the yield of C<sub>6</sub>H<sub>6</sub>, the effluent from step three is fed again to the DHA unit in the subsequent cycle. Brady *et al.*<sup>240</sup> demonstrated each of these above steps. Based on extrapolation, the upper limit of aromatics yield for such a process was estimated to be ~43%.<sup>240</sup> However, an integrated CL-DHA process under such a multi-step scheme has yet to be demonstrated. Furthermore, the oxidation of Fe to Fe<sub>3</sub>O<sub>4</sub> with steam is thermodynamically limited. It is also to be noted that the regeneration of Fe with water instead of air renders the overall process endothermic. The use of water for regeneration by Brady *et al.*<sup>240</sup> may have been (partially) motivated by the lack of H<sub>2</sub> combustion selectivity of Fe<sub>2</sub>O<sub>3</sub>. Although this CL-enhanced DHA scheme is quite innovative, it could be simplified while accounting for the aforementioned design considerations. For this, other SHC redox catalysts with higher H<sub>2</sub> combustion selectivities should be considered in addition to iron oxides.<sup>83,145</sup>

**Direct conversion of CH<sub>4</sub> to CH<sub>3</sub>OH.** Direct conversion of CH<sub>4</sub> to CH<sub>3</sub>OH (and/or formaldehyde) *via* (R50) is thermodynamically feasible. However, despite over a century of research the development of such a process remains a challenge owing to (i) a large energy barrier associated with the activation of the C–H bonds of CH<sub>4</sub>, and (ii) a lower stability of CH<sub>3</sub>OH than CH<sub>4</sub>, resulting in further oxidation of CH<sub>3</sub>OH to CO<sub>2</sub>.<sup>243</sup> Consequently, CH<sub>3</sub>OH is currently produced on a commercial scale from synthesis gas derived from coal or CH<sub>4</sub>.<sup>244</sup>

To date, no heterogeneous catalyst exists that can activate CH<sub>4</sub> according to reaction (R50) while simultaneously inhibiting CH<sub>3</sub>OH oxidation at practical yields.<sup>245</sup> Interestingly, in nature monooxygenase (MMO) enzymes convert CH<sub>4</sub> selectively into CH<sub>3</sub>OH at ambient temperature.<sup>246</sup> The active sites in these enzymes, *i.e.* binuclear iron and copper centers, can be mimicked using zeolite frameworks. However, so far with Fe and Cu-based zeolites a high selectivity for CH<sub>3</sub>OH is only achievable at very low CH<sub>4</sub> conversions (<0.1%).<sup>247,248</sup> The low CH<sub>3</sub>OH yields of the direct aerobic conversion of CH<sub>4</sub> to CH<sub>3</sub>OH process is a thermodynamic feature that cannot be overcome by the choice of catalyst. Nonetheless, this thermodynamic constraint can be circumvented by using a CL approach and soft oxidants such as H<sub>2</sub>O or N<sub>2</sub>O.<sup>75,76,249–251</sup> For example, using a Cu-exchanged mordenite zeolite, Sushkevich *et al.*<sup>252,253</sup> reported an anaerobic CL process based on the reaction (R66) to avoid the over-oxidation of CH<sub>3</sub>OH. In the proposed three-step process, the Cu-exchanged mordenite zeolite was first activated at 400 °C in a flow of either dry O<sub>2</sub> or He, followed by exposure to CH<sub>4</sub> at 200 °C and 7 bar to form methoxy species. Subsequently, the methoxy species were desorbed in the form of CH<sub>3</sub>OH under a flow of a H<sub>2</sub>O/He mixture. Once the desorption step was complete, the second reaction cycle was started by heating the zeolite again to 400 °C under a flow of dry O<sub>2</sub> or He. This process resulted in a normalized CH<sub>3</sub>OH production rate of ~20 μmole CH<sub>3</sub>OH g<sub>catalyst</sub><sup>-1</sup> h<sup>-1</sup> (based on a cycle time of 6.7 h) with a selectivity of 97%, which is considerably higher than the CH<sub>3</sub>OH production rate of 1.81 μmole CH<sub>3</sub>OH g<sub>catalyst</sub><sup>-1</sup> h<sup>-1</sup> reported for a continuous process using O<sub>2</sub> as the oxidant.<sup>254</sup>



## 4.2 Chemical looping for C<sub>2+</sub> valorization

**4.2.1 ODH of light alkanes.** As illustrated in Section 2, the chemical looping strategy has been explored to intensify the production of ethylene,<sup>111</sup> an important commodity chemical with a worldwide production capacity exceeding 150 million tonnes per year.<sup>255</sup> Compared to the well-established, yet energy- and carbon-intensive ethane (and naphtha) cracking processes, the CL-ODH approach (Fig. 12) has the potential to reduce the energy consumption and CO<sub>2</sub> emissions for ethylene production by as much as 87%.<sup>113</sup> The significant process intensification of CL-ODH is enabled by: (i) built-in air separation *via* chemical looping; (ii) higher ethane conversion with autothermal operations *via* the *in situ* oxidation of hydrogen; and (iii) simplified downstream product separation due to the higher ethylene yield and the combustion of hydrogen.<sup>256</sup> We note that extensive research has been conducted for catalytic ethane ODH in the presence of gaseous oxygen,<sup>257–260</sup> but with limited success,<sup>261,262</sup> The key challenges for catalytic ODH resides in the cost of oxygen, safety issues related to the oxygen/ethane co-feed, the lack of catalysts with high activity and selectivity, and the complexity for the removal of oxygenate byproducts. To date, no suitable ODH catalysts have been identified to achieve sufficiently high olefin yields (~70%) such that ODH could replace the commercial steam cracking processes.<sup>263,264</sup> Moreover, most of ODH reactions were carried out at very low ethane partial pressures (<0.1 bar) due to safety concerns. In contrast, CL-ODH avoids direct contact between ethane and oxygen, provides integrated oxygen separation, and has demonstrated superior olefin yields (>70%)<sup>265</sup> at practical ethane partial pressures (≥0.8 bar). In principle, the same CL-ODH strategy can be applied to other light alkanes such as propane and butane. The following discussions provide a brief account of the chemical looping studies related to ethane and propane conversion to light olefins.

The chemical looping operations illustrated in Fig. 12 can be realized by two types of redox catalysts with distinct functions.<sup>266</sup> The Type I redox catalysts function primarily as SHC materials. Using Type I catalysts, ethylene production is realized by a combination of dehydrogenation (DH) or cracking reaction (for olefin and hydrogen production) and SHC reaction (for the

*in situ* oxidation of the hydrogen byproduct, reaction (R16)). In comparison, Type II redox catalysts catalyze ethane ODH reactions while donating their active lattice oxygen.

**Type I: SHC redox catalysts.** While it could be argued that Type II redox catalysts are more desirable due to their multi-functionality, Type I redox catalysts could offer better flexibility since they can be tuned independently for various operating conditions and reactor configurations. For instance, SHC redox catalysts have been explored to enhance both DH and thermal cracking reactions with a temperature range varying from 450 to 900 °C.<sup>82,83</sup>

The use of SHC to enhance catalytic DH reactions (Route a in Fig. 12) was proposed in the late 1990s by Imai *et al.*<sup>267,268</sup> in the context of propane dehydrogenation. While both co-feeding oxygen<sup>269</sup> and redox mode operation using lattice oxygen<sup>269</sup> were investigated, the latter was surmised to be superior based on both experimental data and reactor simulations.<sup>268,270,271</sup> Resulting from their significant potential, various oxides were explored as SHC redox catalysts in the temperature range of 550–800 °C. Representative materials showing high (>90%) SHC selectivities, defined as the conversion of the hydrogen divided by the conversion of all combustible species, include Sb<sub>2</sub>O<sub>4</sub>, In<sub>2</sub>O<sub>3</sub>, WO<sub>3</sub>, Bi<sub>2</sub>O<sub>3</sub>, PbCrO<sub>4</sub>, doped ceria, ion-exchanged ZSM-5, *etc.*<sup>268,269,272–277</sup> Bi, In, Pb, and Sb oxides often suffer from a poor redox stability owing to relatively low melting temperatures of their reduced states. Ceria doped with W, Bi, Cr or Pb, were found to be stable in the temperature range 500–600 °C with selectivities of up to 98% for hydrogen combustion in the presence of C<sub>2</sub> and C<sub>3</sub> hydrocarbons and oxygen storage capacities as high as 2 wt%.<sup>273,278–282</sup> Mn- and La-containing perovskites also showed a high SHC selectivity and stability in the presence of C<sub>3</sub> hydrocarbons at 550 °C.<sup>84</sup> It is noted, however, that the SHC selectivity tends to decrease with increasing temperature due to the lower activation energy for hydrogen combustion compared to hydrocarbon oxidation. Therefore, these early SHC studies focusing mostly on operating temperatures < 650 °C were exclusively geared towards dehydrogenation applications in combination with a DH catalyst such as supported PGM (*e.g.* Pt) and chromium oxides.<sup>259,278,283</sup>

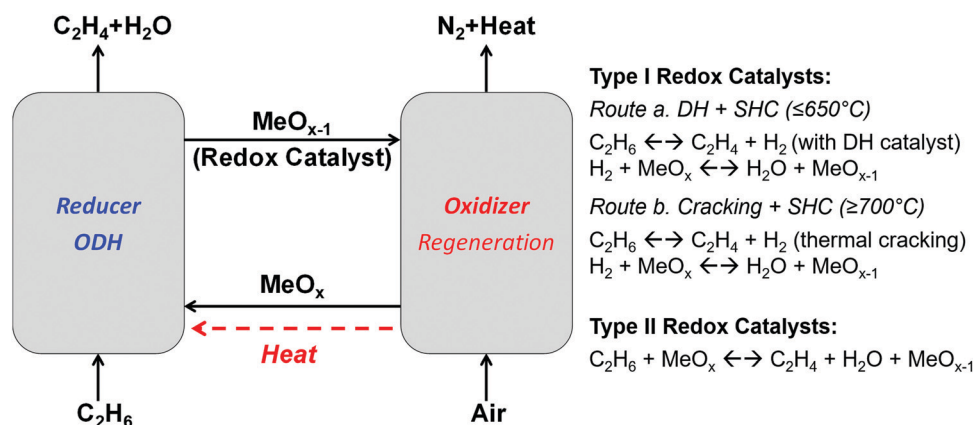


Fig. 12 Schematic of CL-ODH, redox catalyst types, and reaction schemes. Ethane is used as an example for light alkanes for illustration purpose.



Another challenge with the DH and SHC concept resides in the incompatibility of the catalysts and their operating conditions. While physical and chemical interactions between the catalysts themselves can be avoided by spatially separating them (*e.g.* in a packed bed with alternating DH catalyst and SHC redox catalyst layers), active sites and dominating reaction pathways on DH catalysts can be affected by the steam generated from the SHC material. For instance, steam can lead to undesirable reforming reactions on PGM-containing catalysts. For oxide-based DH catalysts, steam (or air during the regeneration step) can affect the oxidation state of the active metal, induce changes in the surface compositions, and hence negatively affecting the DH performance. Except for very few studies,<sup>269</sup> the aforementioned studies used hydrogen and a hydrocarbon mixture (*e.g.* hydrogen, ethane, and ethylene) to evaluate the redox catalysts' SHC performances. The incompatibility issues between DH and SHC have largely been unaddressed to date. Development of DH catalysts that are stable under redox conditions and in the presence of steam is therefore the key to apply the DH + SHC approach for CL-ODH.

SHC redox catalysts can also be integrated with thermal cracking reactions (Route b in Fig. 12). Unlike surface-catalyzed DH reactions, thermal cracking proceeds *via* gas-phase radical reactions and so steam generated by SHC would not negatively affect cracking reactions. Under this approach, the CL-ODH reactions proceed *via* parallel gas-phase and surface pathways, *i.e.* hydrogen generated from gas phase cracking reactions is selectively combusted on the surface of the redox catalysts.<sup>80,269</sup> In some cases, the redox catalyst surface can also function as a radical initiator to enhance the gas phase reactions (*i.e.* surface-initiated homogeneous reactions).<sup>87</sup> It is noted that the initiation of radicals from alkanes typically requires temperatures close to or higher than 700 °C. While such a temperature range is not uncommon for CLC, the inverse relationship between SHC selectivity and reaction temperature for most oxide redox catalysts dictates that many of the SHC catalysts reported for DH + SHC, which are geared towards operating at 650 °C or lower, would not be compatible with thermal cracking reactions. The high operating temperature can also be challenging for oxides or metals with low melting points or high vapor pressures. To address these challenges, Dudek *et al.*<sup>83</sup> investigated a number of Mn-containing redox catalysts including CaMnO<sub>3</sub>, SrMnO<sub>3</sub>, and Mg<sub>6</sub>MnO<sub>8</sub> for selective hydrogen combustion using hydrogen and ethylene mixtures at temperatures up to 850 °C. The effect of promoters was also studied by impregnating Na<sub>2</sub>WO<sub>4</sub> on each of these oxides. In general, Na<sub>2</sub>WO<sub>4</sub> was effective in inhibiting the formation of CO<sub>x</sub>, but negatively affected the redox activity of the mixed oxide. The operating temperature windows, *i.e.* the optimal temperature ranges with a high activity and selectivity towards hydrogen combustion, for the six redox catalysts investigated ranged between 550 and 850 °C. The large and tunable operating temperature window opens up the potential opportunity for integration of a SHC redox catalyst with thermal cracking. The effectiveness of CL-ODH under thermal cracking conditions was confirmed with Na<sub>2</sub>WO<sub>4</sub> promoted Mn–Mg and Mn–Si oxides.<sup>284</sup> Resulting from

their high activity and high temperature SHC selectivity, ethylene yields up to 68% were demonstrated with CO<sub>x</sub> selectivities as low as 1.9%. Negligible coking was observed in these studies. This is not surprising considering the redox catalysts' high activity for lattice oxygen donation, which can actively inhibit coke formation. Na<sub>2</sub>WO<sub>4</sub> was important for obtaining a high olefin selectivity since it enriches on the oxide surface and suppresses the deep oxidation of hydrocarbons.<sup>285</sup> In the case of the Mg<sub>6</sub>MnO<sub>8</sub> system, a mixed oxide with a cation deficient rocksalt structure, near surface Mn<sup>4+</sup> was suppressed by more than 85%.<sup>80,81</sup> Further characterizations by low energy ion scattering (LEIS), *in situ* XRD, and differential scanning calorimetry indicated that the Na<sub>2</sub>WO<sub>4</sub> is likely to form a molten layer on the oxide surface under the operating conditions and thus acts as a physical blocker to suppress surface catalyzed C–H bond activation.<sup>286</sup> The corresponding process analysis further highlighted the advantages of CL-ODH of ethane over steam cracking.<sup>113</sup> These include higher single-pass yields, a reduction in the total energy demand by >80% and a reduction in CO<sub>2</sub> emissions by >80%. Importantly, CL-ODH is self-sufficient from a process heat requirement point of view, making it a highly efficient and low emission technology ideal for distributed systems.

*Type II: ODH redox catalysts.* Many heterogeneous catalysts used in conventional ethane ODH (where oxygen is co-fed with the ethane) are redox-active.<sup>287</sup> In the presence of gaseous oxygen (or, as was shown recently, mild oxidants such as steam),<sup>288</sup> the reaction proceeds through a MvK mechanism<sup>289</sup> in which ethane is activated and dehydrogenated by surface metal-oxide species, thereby reducing the metal oxide and creating oxygen vacancies. The vacancies on the surface are subsequently replenished by gaseous oxygen, which is dissociated and incorporated at surface sites different from the ODH reaction sites.<sup>290–292</sup> The overall reaction is therefore facilitated by the migration of oxygen anions through the lattice and/or on the surface. The most common redox-active heterogeneous catalysts for ODH of light alkanes are based on vanadium oxide<sup>293–295</sup> with different supports, such as alumina,<sup>296–299</sup> alumina modified with zirconia<sup>300,301</sup> or silica.<sup>302–305</sup> Nickel oxides<sup>306,307</sup> and cobalt oxides<sup>308,309</sup> have also been examined.

Since a key characteristic for the MvK mechanism is that the overall reaction can be carried out under a “swing mode”, *i.e.* separating the reduction and oxidation into two sequential steps identical to chemical looping operations, it is not surprising that many of the redox-active oxide catalysts can act as redox catalysts in CL-ODH schemes. For example, Hossain and de Lasa developed a number of cyclically stable VO<sub>x</sub>/γ-alumina-based catalysts for the CL-ODH of ethane in a continuous fluidized bed process consisting of an ODH reactor and a catalyst regenerator.<sup>298,299,310</sup> Ethylene selectivities ranging from 80–90% at temperatures between 550–600 °C were reported. The corresponding ethane conversions, however, were relatively low (<10%).<sup>300,311,312</sup> The same class of catalysts was also used for the CL-ODH of propane in the simulated fluidized bed process and propylene selectivities of >90% were obtained at up to 25% conversion of propane.<sup>313–315</sup> The good performance of the most promising





redox catalyst,  $\text{VO}_x/\text{CaO}-\gamma\text{-Al}_2\text{O}_3$ , was attributed to an intermediate acidity of the catalyst and moderate active site-support interactions.<sup>85</sup> Encouraging results for ethane CL-ODH were also obtained with Mo-based redox catalysts, which gave ethylene selectivities of 93–95% and conversion up to 66.5% at 600 °C.<sup>297,316</sup> The oxygen storage capacity of these Mo-based redox catalysts was relatively low (<0.4 wt%), resulting in a decreasing conversion of ethane with time. Nonetheless, in a continuously operating riser reactor system, an ethane conversion of 12% at ethylene selectivities >90% was obtained with a mixed Mo–Te–V–Nb oxide supported on  $\gamma\text{-Al}_2\text{O}_3$  when the contact time of ethane feed and redox catalysts was 1–2 s at 500–575 °C.<sup>317</sup> The same authors also showed that the performance of Mo-based redox catalysts is strongly affected by the support material ( $\text{Al}_2\text{O}_3$ ,  $\text{Ga}_2\text{O}_3$  or  $\text{Y}_2\text{O}_3$ ), leading to an increased ethane conversion and ethylene selectivity with a decrease in the content of  $\text{MoO}_3$ ; the best results were obtained with 5 wt%  $\text{MoO}_3$  on alumina, giving 90% selectivity for ethylene at an ethane conversion of 30%.<sup>318,319</sup> Novotny *et al.*<sup>284</sup> investigated the effect of impregnating  $\text{MoO}_3$  onto an  $\alpha\text{-Fe}_2\text{O}_3$ -based redox catalyst, which improved the oxygen storage capacity from 0.28 wt% to 0.62 wt%. A  $\text{MoO}_3$  and  $\text{Fe}_2(\text{MoO}_4)_3$  enriched surface layer was formed upon calcination. At 600 °C, the ethylene selectivity was as high as 62% while  $\text{CO}_2$  formation was suppressed when compared to pure  $\alpha\text{-Fe}_2\text{O}_3$ . Supported chromium oxide catalysts are widely used in catalytic dehydrogenation,<sup>320</sup> *e.g.* the commercial CATOFIN<sup>®</sup> process. Chromium oxides were also shown to perform reasonably well in CL-ODH of propane with potential use in a moving bed process.<sup>321–323</sup> The oxygen storage capacity was determined to be crucial for the performance of the redox catalyst, and slightly higher oxygen capacities (up to ~0.5 wt%) resulted in higher yields of propylene (~46% at 630 °C).<sup>324</sup> More recently, Chen *et al.* used to use Mo–V–O mixed oxides as the redox catalyst for CL-ODH of propane to propylene.<sup>325</sup> Attractive propylene yield (89% propylene selectivity at 36% propane conversion) over 100 redox cycles was obtained at 500 °C due to the tailored bulk lattice oxygen *via* atomic-scale doping of Mo in the redox catalyst. Although promising results were obtained from these redox catalysts, the toxicity of Cr and V and thermal stabilities of Mo and V oxides could affect their practical applicability especially in circulating bed systems.

Aside from the redox-active oxides commonly found in heterogeneous DH and ODH catalysts, alkali metal-promoted, Ruddlesden–Popper structured,  $\text{La}_x\text{Sr}_{2-x}\text{FeO}_{4-\delta}$  redox catalyst was recently shown to be active for CL-ODH of ethane.<sup>79,128</sup> Up to 90% ethylene selectivity and 61% conversion were observed at 700 °C, with near 100%  $\text{H}_2\text{O}$  selectivity. An important finding from this study was that electrophilic surface oxygen species (such as  $\text{O}^-$  and  $\text{O}_2^-$ ), which limit the selectivity of the ODH reaction and account for the deep oxidation of the alkanes and olefins, can be suppressed by enrichment of alkali metal oxide on the surface of the redox catalyst. A thin layer of  $\text{Li}_2\text{O}$  on the surface likely served as a barrier to inhibit the outward diffusion of  $\text{O}^{2-}$  and its evolution into electrophilic oxygen species on the surface.

A common challenge with these Type II ODH redox catalysts is the lack of oxygen storage capacity. Most of the redox

catalysts reported to date exhibit <1 wt%, and in many cases <0.5 wt%, oxygen storage capacity. As noted above, the utilization of lattice oxygen in CL-ODH is significantly more efficient than that in CLC. For instance, combusting one mole of propane under CLC consumes ten moles of lattice oxygen whereas under CL-ODH it consumes only one. The order of magnitude increase in oxygen utilization efficiency make these low capacity redox catalysts potentially feasible for industrial applications, provided that high olefin selectivity, redox activity, and stability are demonstrated. Another challenge is the trade-off between conversion and selectivity especially in many of the V- and Mo-based redox catalysts, *i.e.* higher alkane conversion (*e.g.* by increasing the residence time) often leads to a significantly decreased olefin selectivity. This could result from the re-adsorption of the olefin products and/or the presence of distinct active sites for selective and non-selective reactions. In-depth understanding of the active sites and reaction pathways would be crucial for rational optimization of the redox catalysts. Although a large body of literature is available on conventional, catalytic ODH systems, the exact mechanisms for many redox-active ODH catalysts are still under debate.<sup>259,326</sup> The dynamic nature of redox catalysts undergoing chemical looping reactions also makes mechanistic investigation on CL-ODH even more challenging than catalytic ODH. Therefore, further investigation on the mechanistic aspects of the redox catalysts is an important, yet challenging task. Although we did not separately discuss ethane and propane CL-ODH in this section, we note that propane (and  $\text{C}_{4+}$ ) is significantly less stable than ethane. The tendency for C–C bond cleavage makes it more difficult to develop highly selective redox catalysts for CL-ODH of propane and  $\text{C}_{4+}$  to olefins. Thus, a redox catalyst optimized for CL-ODH of ethane may not automatically be applicable for propane and  $\text{C}_{4+}$  conversion.

**4.2.2 Epoxidation of olefins.** The epoxidation of olefins (R51) is another example for oxidation reactions where yields are generally low due to the non-selective oxidation of the educts and the epoxides; yet, these catalytic reactions are of enormous relevance for the chemical industry.<sup>327</sup>

So far, only the epoxidation of ethylene has been realized industrially using molecular oxygen as the sole oxidant.<sup>328</sup> The selectivity to ethylene oxide is typically in the range of 85–90% at ~10% ethylene conversion with optimized silver-based catalysts.<sup>329–331</sup> A chemical looping approach was recently explored to undertake the epoxidation reaction by substituting co-fed molecular oxygen with lattice oxygen (Fig. 13).<sup>78</sup> Here, Ag (15 wt%) was impregnated onto the perovskite  $\text{SrFeO}_3$  that possesses an oxygen storage capacity of up to ~2 wt%. The main rationale for selecting  $\text{SrFeO}_3$  was its excellent ability to spontaneously release  $\text{O}_2$  at relatively low temperatures.<sup>136</sup> The authors proposed that oxygen was transported from subsurface  $\text{SrFeO}_3$  to the surface of the redox catalyst and thus formed oxygen adatoms on the silver surface. The vital role of atomic oxygen adsorbed on the silver surface for catalyzing the oxidation of ethylene has been discussed extensively in the literature,<sup>331,332</sup> as well as the availability of subsurface oxygen to achieve a high ethylene oxide selectivity.<sup>333</sup> Furthermore, it was found in



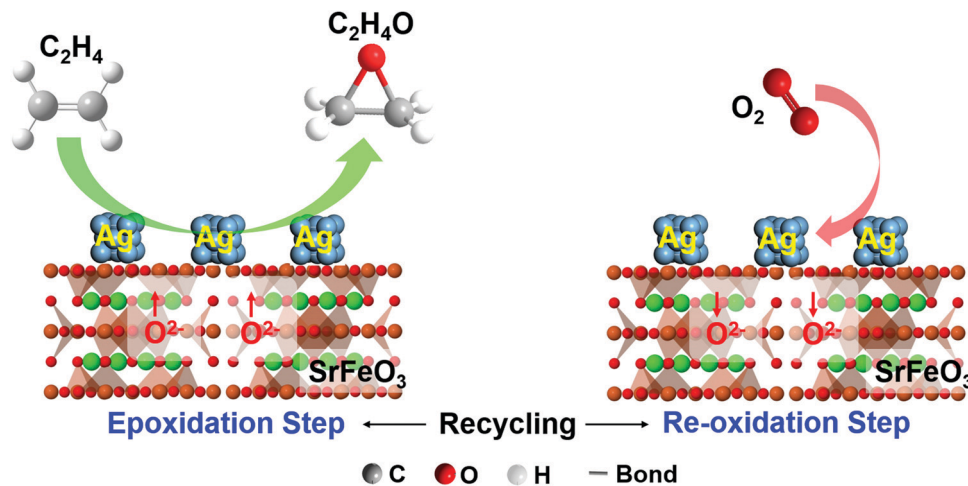


Fig. 13 The reaction mechanism of chemical looping epoxidation using a silver-modified  $\text{SrFeO}_{3-x}$  redox catalyst.

previous studies that higher selectivities could be obtained with low surface area supports,<sup>334,335</sup> a characteristic of most redox catalysts used in chemical looping applications. The selectivities and conversions reported by Chan *et al.*<sup>78</sup> were, however, very low (the ethylene oxide selectivity was <20% and the ethylene conversion was <4% at 270 °C). Previous work showed that the dispersion of Ag, the morphology and temperature treatment of the support all affect, besides operational parameters, the performance of the catalyst.<sup>336–338</sup> It is also possible that the exposed surface of  $\text{SrFeO}_3$  is largely responsible for non-selective oxidation reactions and the activity of the oxide substrate to catalyze non-selective oxidation reactions should not be overlooked. Another challenge observed by Chan *et al.* was the low rate of oxidation when the redox catalyst was regenerated in air, causing a gradual loss in oxygen capacity.<sup>78</sup> A similar problem was reported for vanadium phosphorous oxide redox catalysts, as will be discussed in Section 4.2.3. The problem was overcome by modifying the  $\text{SrFeO}_3$  with 5 mol%  $\text{CeO}_2$ , which also improved the cyclic stability and ultimately gave 60% selectivity towards ethylene oxide at 10% ethylene conversion.<sup>339</sup> The authors proposed that  $\text{CeO}_2$  may have acted as oxygen gateway (see Section 3.2.2) and thus improved the re-oxidation of the redox catalyst. The challenge for further improvement will thus be to find oxygen storage materials with sufficient reduction/regeneration activities at temperatures <300 °C while suppressing over-oxidation of ethylene. Despite the relatively low reaction temperatures of 200–280 °C typically employed for the epoxidation of ethylene, Ag-based catalysts tend to deactivate owing to the sintering of the silver particles,<sup>340–342</sup> and this may be even more problematic when the redox catalysts are regenerated in an exothermic oxidation reaction.

In principle, the chemical looping epoxidation approach can certainly also be applied to the selective oxidation of propylene, where it was shown recently that mixed metal oxides are catalytically active in a conventional oxygen co-fed mode. For a mixed  $\text{MoO}_3/\text{Bi}_2\text{SiO}_5$  catalysts supported on  $\text{SiO}_2$ , propylene oxide selectivities of 55% were reported at a propylene conversion of 22% at 400 °C for a diluted feed with a molar ratio of  $\text{O}_2/\text{C}_3\text{H}_6$  of 4:1,<sup>343</sup> whereas high formation rates of propylene

oxide were observed for a  $\text{RuO}_2/\text{CuO}$  catalyst supported on  $\text{SiO}_2$ .<sup>344</sup> Using a copper–manganese mixed metal oxide catalyst, Seubsai *et al.*<sup>345</sup> found selectivities >30% towards propylene oxide at up to 1.5% propylene conversion at 300 °C. These catalyst systems are structurally and chemically similar to typical chemical looping redox catalysts, indicating the potential for future work in this field.

**4.2.3 Maleic anhydride production.** ODH in a redox mode was demonstrated commercially by DuPont for the oxidation of *n*-butane to maleic anhydride to produce maleic acid (R53).<sup>86,346</sup> Up to 6500 kg  $\text{h}^{-1}$  of maleic acid were produced using a vanadium phosphorous oxide (VPO) redox catalyst in a circulating fluidized bed reactor operated at ~400 °C.<sup>347</sup> The process was developed over a period of more than ten years, as summarized by Contractor, also highlighting the challenges associated with redox catalyst and reactor design.<sup>348,349</sup> Selectivity of near 80% was claimed at a conversion of 40%.<sup>327</sup> Nevertheless, the plant was shut down in the early 2000s, partially owing to economic reasons related to a rapidly changing market situation for different raw materials and intermediates,<sup>350</sup> but also owing to problems with catalyst loss through attrition<sup>351</sup> and deteriorating oxidation kinetics of the VPO catalyst, which resulted in a gradual loss in oxygen capacity during operation such that  $\text{O}_2$  had to be co-fed.<sup>352</sup> We also note that the oxygen capacity of the VPO redox catalyst was rather low to begin with, corresponding to merely a few monolayers of lattice oxygen near the catalyst surface.<sup>353,354</sup>

### 4.3 Chemical looping involving liquid and solid fuels

**4.3.1 Liquid fuels driven processes.** The CLBC approach can readily be extended to liquid hydrocarbons. In fact, it has been successfully employed to reform various liquid fuels, such as alcohols (*e.g.* methanol or ethanol),<sup>355–362</sup> glycerol,<sup>363</sup> waste cooking oil,<sup>364</sup> liquid alkanes (*e.g.* dodecane),<sup>365</sup> kerosene,<sup>366</sup> heavy oil,<sup>365</sup> and biomass tar.<sup>367–370</sup> In addition to reforming, liquid fuels can also decompose or completely oxidize in the fuel reactor. For example, Ochoa *et al.*<sup>359</sup> explored the nature of the surface species and gaseous products formed during



reduction of different spinel mixed oxides using ethanol. It was found that ethanol always first dehydrogenates to acetaldehyde on the surface of the spinel. Subsequently, depending upon the spinel composition, acetaldehyde (i) partially oxidizes to acetates (Ni-based spinel), (ii) decomposes to CO and CH<sub>4</sub> (Co-based spinel), or (iii) completely oxidizes to CO<sub>2</sub> and H<sub>2</sub>O (Fe-based spinel). The design criteria of redox catalysts and operating conditions for liquid fuels driven chemical looping processes are similar to those of the gaseous hydrocarbon-based chemical looping processes. However, special care must be taken when using viscous liquids such as tar or heavy oil to avoid their condensation upstream and/or downstream of the fuel reactor. Similar to Type II ODH process, the chemical looping approach has been applied for the oxidative cracking of naphtha. Process simulations indicate that over 50% energy and CO<sub>2</sub> savings can be achieved *via* the chemical looping route.<sup>112</sup> Using perovskite-based redox catalysts, Dudek *et al.*<sup>87</sup> demonstrated close to 70% olefin selectivity with a CO<sub>x</sub> selectivity as low as 0.3%.

**4.3.2 Solid fuels driven processes.** Owing to the slow rates of solid-solid reactions, using solid fuels directly for CLBC is challenging. Therefore, solid fuels (such as biomass or coal) are typically gasified in a separate unit and the resulting synthesis gas is fed to the chemical looping process designed for gaseous fuels.<sup>371–374</sup> To avoid dilution of the synthesis gas with N<sub>2</sub>, the gasification reaction needs to be carried out with pure O<sub>2</sub> or a mixture of O<sub>2</sub> and steam/CO<sub>2</sub>. Therefore, an air separation unit would be required. Several strategies to extend CLBC to solid fuels are summarized below:

(i) The solid fuel can be gasified *in situ* in the presence of a redox catalyst using steam or CO<sub>2</sub> as the fluidizing gas. This strategy not only eliminates the need for a separate gasification unit but also enhances the rate of gasification.<sup>375,376</sup> However, owing to the inherently slow nature of the gasification reaction, it is difficult to achieve complete gasification of the solid fuel *in situ*.

(ii) Depending upon their thermodynamic characteristics, certain oxygen carriers can undergo spontaneous decomposition at high temperatures to release molecular oxygen.<sup>377,378</sup> The use of such oxygen carriers allows the direct combustion of a solid fuel *via* a gas–solid reaction, *i.e.* without an intermediate gasification step. This process configuration, typically referred to as chemical looping with oxygen uncoupling (CLOU), significantly increases the rate of fuel oxidation, thereby allowing the complete combustion of the solid fuel. Furthermore, instead of fully oxidizing the solid fuel it is also possible to gasify it by controlling the partial pressure of oxygen released by the redox catalyst.<sup>379</sup>

(iii) Recently, it has been proposed to use molten oxygen carriers or reaction system for CL-methane POx, CLC, and CL separation to circumvent the challenges associated with the solid-based chemical looping systems, such as carbon deposition, thermal sintering and low reaction rates.<sup>380</sup> Wang *et al.*<sup>381</sup> proposed a molten slat alkali carbonate based CL-methane POx system for the coproduction of synthesis gas and metallic Zn. The molten reaction system enables to intensify heat and mass transfer during the cyclic redox, increasing the energy

utilization efficiency. Sarafraz *et al.*<sup>382–384</sup> investigated the thermodynamic feasibility of using liquid oxygen carriers for the partial oxidation of fuel to synthesis gas and identified Cu, Pb and Sb as the most promising candidates. Upham *et al.*<sup>385</sup> demonstrated a molten bromide-oxide based chemical looping for the reactive separation of HBr in a halogen-based natural gas conversion. Nickel-alkali bromide and oxide were switched in molten reactor for the conversion of HBr to Br through the MeBr/MeO redox pair. Although the use of molten redox catalysts avoids the aforementioned issues and enhances the rate of reaction, it also introduces new challenges such as evaporation or solidification of liquid oxygen carriers, continuous circulation of liquid metals/oxides, large pressure drops across the reactors, *etc.*

(iv) For solid fuels with high volatile contents, *e.g.* biomass or low-rank coal, it is possible that a redox catalyst can be used to generate value-added products *via* pyrolysis-type of processes – an area that has not been investigated extensively by the chemical looping community.<sup>44,386</sup> It is noted that pyrolysis products with less oxygenates are typically desired, which may conflict with typical chemical looping operations.

#### 4.4 Synergy between oxide and calcium/carbonate looping

CaO-Based sorbents, often in the form of naturally-occurring lime, are used widely in applications such as flue gas desulfurization<sup>387–389</sup> and sorption-enhanced steam methane reforming (SE-SMR) to produce H<sub>2</sub>.<sup>390–393</sup> In SE-SMR, CO<sub>2</sub> is removed continuously in the WGS reaction step, thus shifting the equilibrium to the product side according Le Chatelier's principle.<sup>394</sup> The H<sub>2</sub> yield can thereby be increased to ~97 vol% at 550 °C.<sup>395</sup> The WGS reaction is relevant also in gasification reactions, and generally in schemes where mixtures of CO, CO<sub>2</sub>, H<sub>2</sub> and H<sub>2</sub>O co-exist at high temperature, and so sorbents for CO<sub>2</sub> are conventionally used.<sup>396</sup> Lately, CaO-based sorbents have also gained considerable attention for carbonate looping (CaL),<sup>397–405</sup> a post-combustion CO<sub>2</sub> capture technique.<sup>402,406</sup> A key advantage of using CaO for the sorption of CO<sub>2</sub> is its high theoretical uptake of 0.78 g CO<sub>2</sub> per g CaO, according to reaction (R68) in Table 2. The carbonation reaction is exothermic whereas the reverse reaction, known as the calcination reaction, is endothermic, giving a stream of concentrated CO<sub>2</sub>.<sup>407</sup> Carbonate and oxide (chemical) looping processes can be combined to improve efficiencies of both processes, as discussed below.

##### 4.4.1 Reaction schemes

*Chemical looping with sorbent enhancement for heat-neutral sorbent regeneration.* Besides permitting higher yields of H<sub>2</sub> in the SE-SMR/gasification scheme, the exothermic carbonation reaction can provide a large amount of heat for the strongly endothermic methane reforming or the gasification reaction. Meanwhile, the regeneration of the sorbent *via* the highly endothermic calcination reaction is critical and requires high reaction temperatures (> 750 °C). Performing the calcination reaction in the presence of an exothermic reaction would balance the heat requirements. Such exothermic reaction could be the oxidation of a reduced oxygen carrier, as suggested by Lyon and Cole,<sup>163,408</sup> *e.g.* the simultaneous oxidation of the



Ni-based reforming catalyst. However, a relatively high ratio of Ni to  $\text{CaCO}_3$  ( $>0.75$ ) would be required to balance the heats of reaction, resulting in an increase in the catalyst cost, and the  $\text{CO}_2$  produced is diluted with  $\text{N}_2$  when air is used for the re-oxidation of Ni. In addition, the oxidation of the reforming catalyst at high temperature is likely to promote its sintering and deactivation.

#### Chemical looping with sorbent enhancement for $\text{CO}_2$ capture.

The simultaneous oxidation and calcination reactions eliminate the benefit of CaO-based sorbents to produce a pure stream of  $\text{CO}_2$ . In CaO-based  $\text{CO}_2$  capture architectures, the calcination reaction can be performed in a reactor operated in an oxy-fuel mode, where the heat is provided by combusting additional fuel (e.g. natural gas) in an atmosphere containing  $\text{O}_2$  diluted with  $\text{CO}_2$ .<sup>409–411</sup> Replacing molecular oxygen with an oxygen carrier that reduces in an exothermic reaction while fully combusting the fuel would enable both the balancing of the endothermicity of the calcination reaction and generating a pure stream of  $\text{CO}_2$ .<sup>412</sup> A few redox pairs, e.g.  $\text{CuO}/\text{Cu}$  or  $\text{Mn}_2\text{O}_3/\text{Mn}_3\text{O}_4$ , can be reduced exothermally with  $\text{CH}_4$  and other hydrocarbons.<sup>39</sup> This feature can be applied to integrate the heat released from the CLC reduction step with the endothermic calcination step to produce concentrated  $\text{CO}_2$ . A reaction scheme where CuO-based chemical looping combustion is combined with SE-SMR is shown in Fig. 14a and at least three reactors are required to obtain separate streams of  $\text{H}_2$  and  $\text{CO}_2$ .<sup>413</sup>

For carbonate looping as a  $\text{CO}_2$  capture technique, heat integration is vital to be economically competitive with other  $\text{CO}_2$  capture techniques. In analogy to the SE-SMR process, CuO-based oxygen carriers have mostly been investigated to render the heat exothermic during the exothermic calcination step.<sup>414–417</sup>

Unlike the CLC reduction step that is often endothermic, the regeneration step is always exothermic for monometallic oxides. Therefore, integration of the heat released in the CLC regeneration step with the sorbent calcination step can also

result in concentrated  $\text{CO}_2$  if a heat transfer agent is used to avoid direct contact between the air (in the regeneration step and the oxidizer) and  $\text{CO}_2$  (from calcination in the calciner). Both ALSTOM and General Electric proposed and investigated such schemes (details on these two processes have been summarized by Fan<sup>122</sup>). In ALSTOM's Hybrid Combustion–Gasification Process for hydrogen generation, a  $\text{CaSO}_4$  oxygen carrier is employed to generate the heat for the calcination of the CaO sorbent. Instead of oxidizing the oxygen carrier in the vicinity of the  $\text{CaCO}_3$ , bauxite is used to transfer the heat from the oxidation reactor to the calcination reactor. Similarly, General Electric's Fuel-Flexible Advanced Gasification–Combustion Process also adopted CLC reactions to provide the heat for calcination. In this case, an iron oxide-based oxygen carrier was used as the heat transfer agent.

*Sorption-enhanced methane reforming coupled with chemical looping conversion schemes.* The thermodynamic properties of some oxygen carriers permit the splitting of  $\text{CO}_2$  or  $\text{H}_2\text{O}$  to produce CO or  $\text{H}_2$  upon oxidation (see Section 4.1.1). Since the rate of reduction of many oxygen carriers, e.g.  $\text{Fe}_2\text{O}_3$ , with primary fuels such as  $\text{CH}_4$  is low, it was proposed to convert  $\text{CH}_4$  first to a synthesis gas over a catalysts (e.g. Ni; in analogy to the dry reforming of methane (DRM)) and use the synthesis gas to reduce the oxygen carrier.<sup>94,418</sup> Depending on the oxidant used ( $\text{CO}_2$  or  $\text{H}_2\text{O}$ ) for the subsequent regeneration of the oxygen carrier, CO or  $\text{H}_2$  can readily be produced.

This reaction concept can be extended by adding a CaO-based sorbent (Fig. 14b).<sup>419,420</sup> Analogously to SE-SMR, the heat of sorption from the carbonation reaction (reaction (R68)) balances the endothermic reforming and reduction reactions. In the following oxidation step,  $\text{CO}_2$  is released from the sorbent and oxidizes the oxygen carrier while producing CO. The chemical looping concept, where in theory three moles of  $\text{CO}_2$  per mole  $\text{CH}_4$  are converted to four moles of CO, has been termed super-dry reforming of methane (R67). Importantly, in such a process the formation of an equilibrium mixture of  $\text{CO}_2$ , CO,  $\text{H}_2\text{O}$  and  $\text{H}_2$  is circumvented during the



Fig. 14 (a) Reaction scheme with three reactors where CuO-based chemical looping combustion is integrated into the SE-SMR process. (b) Reaction scheme of the super-dry reforming of methane with two reactors, in which the reforming catalyst (Ni) is spatially separated from the oxygen carrier ( $\text{Fe}_3\text{O}_4$ ) and the  $\text{CO}_2$  sorbent.





reforming step, thus avoiding a loss in CO yield through the RWGS reaction (R71). So far, the process has been demonstrated by mixing different materials, *viz.* CaO supported on Al<sub>2</sub>O<sub>3</sub> as CO<sub>2</sub> sorbent, Fe<sub>2</sub>O<sub>3</sub> supported on MgAl<sub>2</sub>O<sub>4</sub> as oxygen carrier, and NiO supported on MgAl<sub>2</sub>O<sub>4</sub> as the CH<sub>4</sub> reforming catalyst, rather than multifunctional redox catalysts. While the theoretical advantages over conventional dry-reforming of CH<sub>4</sub> are apparent, there is still scope for improvement (besides improving the materials used). For example, it was proposed to run the process isothermally. Equilibrium thermodynamics dictate that the release of CO<sub>2</sub> from the CO<sub>2</sub> sorbent requires very low CO<sub>2</sub> partial pressures and for that, an inert gas would likely be needed in the oxidation step (where Fe is oxidized by the CO<sub>2</sub> released from the sorbent). This will inevitably dilute the product gas CO. The product gas CO is diluted further with CO<sub>2</sub> because the conversion of CO<sub>2</sub> to CO over Fe is thermodynamically limited and the ratio of CO/CO<sub>2</sub> in the product gas will decrease further as the Fe-based oxygen carrier is oxidized *via* FeO to Fe<sub>3</sub>O<sub>4</sub>.

**4.4.2 Material design.** The performance and function of CaO-based sorbents can be further improved by doping with active oxides,<sup>414,421,422</sup> addition of supports,<sup>423</sup> acid modifications,<sup>424,425</sup> and morphology control.<sup>421,426</sup> One of the most studied combined oxide and carbonate looping schemes is CuO-based chemical looping combustion coupled with SE-SMR.<sup>427–430</sup> In this process, at least three materials are

required: A catalyst for the reforming of CH<sub>4</sub>, a sorbent for CO<sub>2</sub>, and CuO as the oxygen carrier that can be reduced by various fuels in an exothermic reaction. If CH<sub>4</sub> is used as the fuel, the Cu/Ca ratio needs to be >3.1 to achieve thermally neutral conditions in the calcination step based on the enthalpies of reaction at 900 °C. However, this figure varies somewhat and is subject to the process design, the type of fuel (*e.g.* using CO as the fuel, the ratio would be 1.3) and the heat management strategy.<sup>431,432</sup>

Three active components make the design of suitable, “tri-functional” materials rather challenging;<sup>433</sup> current research thus focuses largely on the development and optimization of separate particle systems<sup>434–436</sup> or bi-functional Ca–Cu<sup>422,437–440</sup> or Ca–Ni<sup>441–443</sup> composites and uses commercially available Ni-based catalyst particles.<sup>444,445</sup> The CO<sub>2</sub> capture capacity of the CaO-based sorbents is linked strongly with the available surface area, which is known to decay due to sintering when the sorbents undergo cycling of carbonation and calcination.<sup>446–448</sup> Producing stable particles with a porous CaO structure in the vicinity of low surface area CuO in a Ca–Cu composite material requires appropriate support materials, *e.g.* MgO<sup>449</sup> or Ca<sub>12</sub>Al<sub>14</sub>O<sub>33</sub>.<sup>450</sup> Recent works<sup>405,414,451</sup> at ETH Zürich have shown how such support materials function as structural stabilizers at the microscopic level and how subtle phase transitions affect the cyclic performance of the CaO-sorbents (Fig. 15). Another recent study

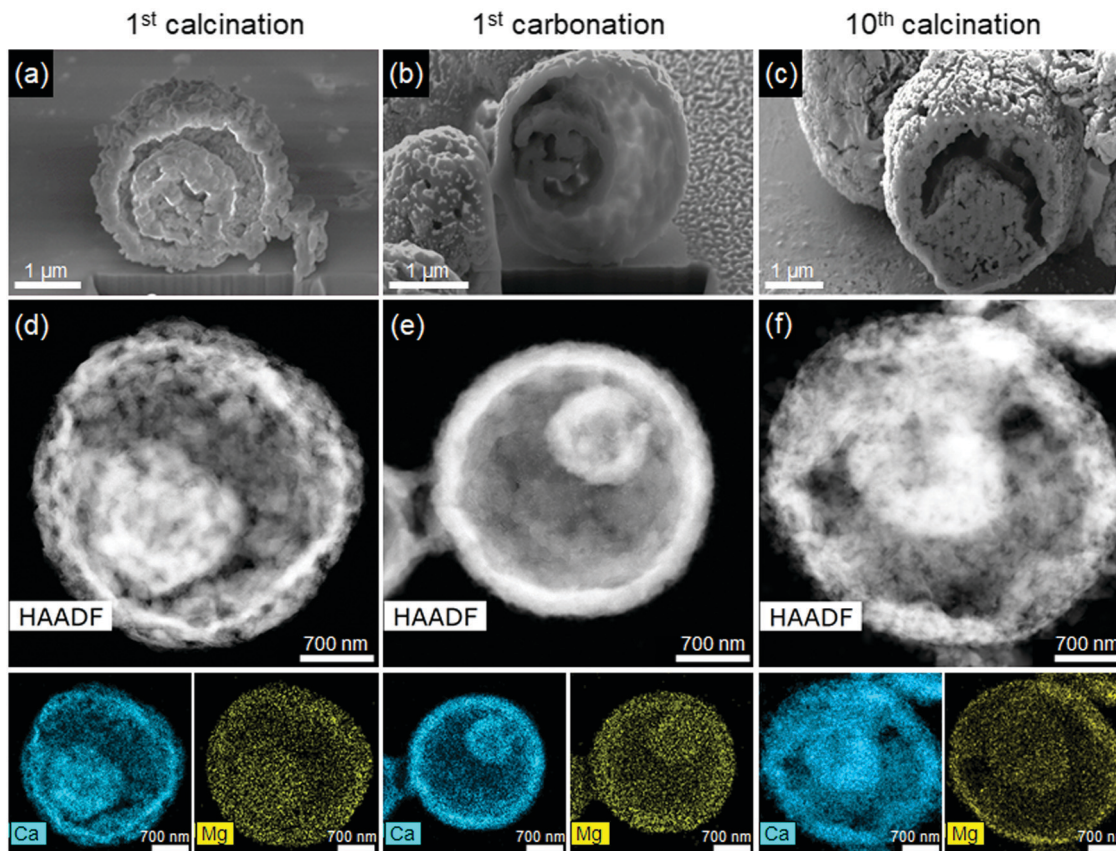


Fig. 15 Morphological changes during cyclic operation of a Ca–Mg mixed oxide-based sorbent. (a–c) SEM images of FIB cross-sections. (d–f) HAADF-STEM images with EDX maps of Ca and Mg at different stages of CO<sub>2</sub> capture/regeneration cycles. Scale bars: 1 μm for (a–c) and 700 nm for (d–f).<sup>451</sup>



reported an interesting phase transition based approach to effectively enhance sorbent stability for the hydrogen production from bio-glycerol.<sup>452</sup>

Conventional design strategies aim to limit the mobility of active species and their growth by using inert support materials that act as physical barriers.<sup>199</sup> New approaches have shown that the active species in a sorbent or redox catalyst may well be mobile, but their self-diffusion is limited if the simultaneous formation of a new phase with the active species is thermodynamically more preferred.<sup>453</sup> For example, Dang *et al.*<sup>452</sup> demonstrated that calcium cobalt, a layered bi-functional material acting as both sorbent and catalyst for the sorbent enhanced reforming of glycerol could effectively be stabilized because the components CaCO<sub>3</sub> and CoO homogenized into calcium cobalt on an atomic level in every calcination stage of the cyclic process instead of remaining segregated and sintering. Well-ordered structures, such as layered oxides, spinels or perovskites are particularly suitable for such approaches.<sup>452,454</sup>

For reaction schemes such as the super-dry reforming of methane, catalyst and oxygen storage material are usually separated spatially.<sup>455</sup> Work by Hosseini *et al.* demonstrated that both the reforming of methane through the catalyst and the reduction of the oxygen storage material can be realized by the same redox catalyst particle provided it is designed carefully in a core-shell structure.<sup>456</sup>

## 5. Summary and perspective

The chemical and petrochemical industry generates an annual revenue in excess of \$4 trillion while consuming 40.9 quadrillion BTU of energy and emitting 2.5 Gigatons of CO<sub>2</sub> (2014 numbers provided by IEA).<sup>457</sup> The significant energy intensity and CO<sub>2</sub> emissions associated with this important industrial sector offer excellent opportunities for process intensification, energy conservation, emission reduction, as well as profitability enhancement. We believe that CLBC represents a promising new frontier in chemical reaction engineering, catalysis, and particle technology to address the aforementioned challenges and opportunities. As discussed in Section 2, the unique advantages of CLBC largely reside in its ability to significantly improve a chemical (and energy) conversion process from a thermodynamic second law standpoint: a strategically designed CLBC scheme, when coupled with tailored oxygen carrier or redox catalyst, can significantly reduce process exergy loss and/or facilitate chemical transitions that would otherwise be impossible under a conventional scheme. From an economic standpoint, reactive separation enabled by oxygen carriers in CLBC offers excellent potential to simplify conventional processes and to reduce the number of unit operations. With the continued technical progress in CLC, a promising carbon capture technology that addresses the carbon intensive, \$1.4 trillion power generation sector, we argue that chemical looping beyond combustion offers equally and perhaps even more exciting opportunities due to their potential for more

efficient lattice oxygen utilization, higher product value, and relative ease for scale-up compared to full scale utility plants.

Although CLBC has attracted increasing attentions over the past 5–10 years, we note this emerging research area is far from being adequately explored particularly considering its significant complexity and excellent potential. From scientific and engineering standpoints, CLBC represents an “ultimate” multidisciplinary and multiscale challenge that requires expertise in various areas including, but not limited to, chemical looping, heterogeneous catalysis, material science, particle technology, reaction and reactor engineering, and process and system engineering. This is due to the uniquely intertwined nature of CLBC. For instance, redox catalyst performance cannot be treated in an isolated manner since it is significantly affected by reactor design and gas–solids contacting patterns as well as the multi-step redox scheme configuration. Meanwhile, the redox catalyst can undergo significant changes in its surface and bulk properties within a CLBC cycle that exceeds complexities typically observed in heterogeneous catalysts. Such complexities, while daunting, are closely associated with the unique advantages CLBC has to offer. They can and should be addressed through continued research *via* interdisciplinary efforts. From a practical standpoint, redox catalysts with high stability and satisfactory long-term performance are critical for CLBC. Moreover, reactor designs that facilitate suitable redox catalyst particle circulation, gas–solids contacting patterns, residence time distributions, and heat integration between multiple reactors are important to enable these promising CLBC technologies.

This article aims to cover all the CLBC schemes to the best of our knowledge. Although a number of promising approaches such as oxidative dehydrogenation of light alkanes and alkene epoxidation are under investigation, significantly greater opportunities still exist both within and beyond these CLBC topics. From a reaction chemistry standpoint, the chemical looping approach is analogous to the MvK mechanism commonly encountered in catalytic oxidation reactions. As illustrated in Fig. 16, CLBC carries out catalytic oxidation cycles *via* a step-wise MvK with macroscopic spatial or temporal separation of the lattice oxygen removal and replenishment events. This is in contrast with conventional MvK, where the catalytic cycle is completed on the surface of an oxide catalyst under steady-state operations. The step-wise operations in CLBC offers both significant advantages (reactive separation, process intensification, and energy/cost reduction) and potential challenges (surface and/or phase changes of the redox catalytic under redox cycles). Considering their similarities, it would not be unreasonable to anticipate catalytic reactions operated under MvK mechanism to be potentially suitable candidates for CLBC. Table 3 summarizes a number of such candidate reactions. Compared to the limited CLBC reactions investigated to date, a significantly number of potential reactions are yet to be explored. In addition, the regeneration step in CLBC, which was demonstrated to be effective for H<sub>2</sub> and CO production *via* CO<sub>2</sub>/H<sub>2</sub>O splitting, could be investigated for hydrogenation or carbonylation reactions. Carriers other than oxygen, *e.g.* nitrogen



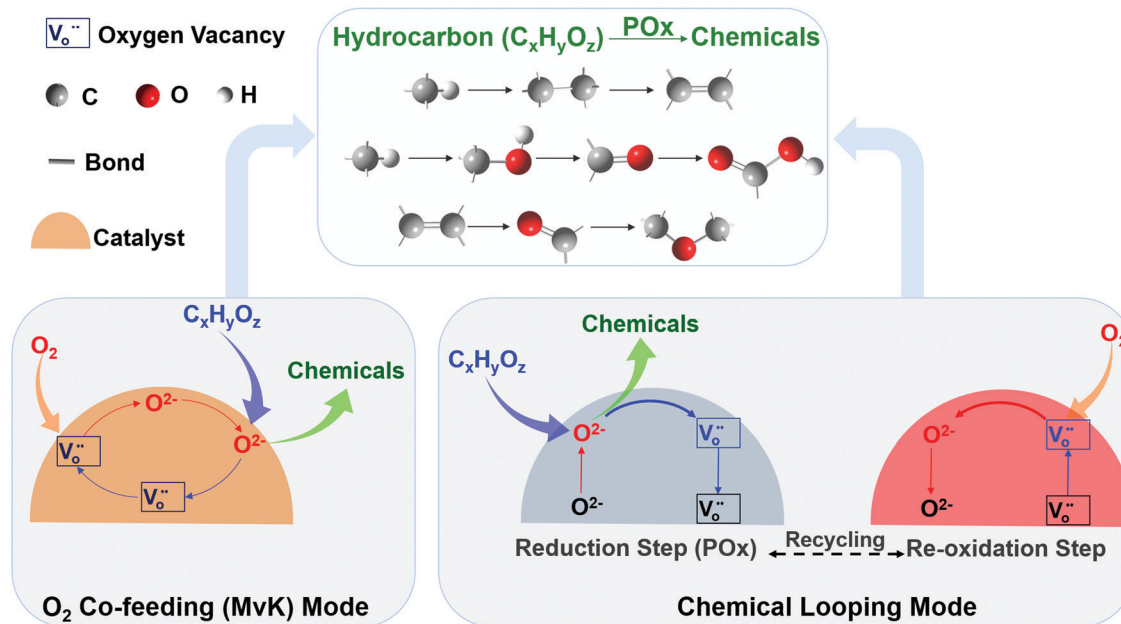


Fig. 16 Comparison of  $O_2$  co-feeding (MvK) and chemical looping modes for partial oxidations and their applications on conversions of hydrocarbon to chemicals (transformation between alkane, alkene, alcohol, aldehyde, ketone, acid, and aromatics).

Table 3 Partial oxidation reactions facilitated by oxide catalysts via the Mars van Krevelen mechanism. The overall oxidation reactions are numbered from (R73) to (R111) and these reactions can potentially be realized by CLBC

Feedstock	Products	Reaction	Heterogeneous catalysts
Methane	Formaldehyde	(R73) $CH_4 + 2CO_2 \rightarrow CH_2O + 2CO + H_2O$	$V_2O_5/SiO_2$ <sup>458</sup>
Methanol	Formaldehyde	(R74) $CH_3OH + 1/2O_2 \rightarrow CH_2O + H_2O$	$Fe_2(MoO_4)_3$ <sup>459,460</sup>
	Methyl formate	(R75) $2CH_3OH + O_2 \rightarrow HCOOCH_3 + 2H_2O$	$V_2O_5/TiO_2$ <sup>461</sup> $SnO_2-MoO_3$ <sup>462</sup>
Ethanol	Acetic acid	(R76) $C_2H_5OH + O_2 \rightarrow CH_3COOH + H_2O$	$V_2O_5/TiO_2$ <sup>463</sup>
	Acetaldehyde	(R77) $C_2H_5OH + 1/2O_2 \rightarrow CH_3CHO + H_2O$	
1,2-Ethanediol	Ethanedial	(R78) $C_2H_6O_2 + O_2 \rightarrow C_2H_2O_2 + 2H_2O$	P-Doped Fe–Mo mixed oxide <sup>464</sup>
Propane	Acrolein	(R79) $C_3H_8 + 3/2O_2 \rightarrow C_3H_4O + 2H_2O$	$NiMoO_4/MoO_3/Te_2MoO_7/P_2O_5$ <sup>465</sup>
	Acrylic acid	(R80) $C_3H_8 + 2O_2 \rightarrow C_3H_4O_2 + 2H_2O$	$NiMoO_4/MoO_3/Te_2MoO_7/P_2O_5$ <sup>465</sup>
Propene	Acrolein	(R81) $C_3H_6 + O_2 \rightarrow C_3H_4O + H_2O$	Fe–Sb mixed oxide, <sup>466</sup> $V_2O_5/Nb_2O_5$ <sup>467</sup>
	Acrylonitrile	(R82) $C_3H_6 + NH_3 + 3/2O_2 \rightarrow C_3H_3N + 3H_2O$	Fe–Sb mixed oxide <sup>466</sup>
	1,5-Hexadiene	(R83) $2C_3H_6 + 1/2O_2 \rightarrow C_6H_{10} + H_2O$	$Bi_2M_{0.1}V_{0.9}O_{5.35}$ (M = Cu or Co) <sup>468</sup>
	Acrylic acid	(R84) $C_3H_6 + 3/2O_2 \rightarrow C_3H_4O_2 + H_2O$	$As_2O_3-Nb_2O_5-MoO_3$ <sup>469</sup>
Butane	1-Butene	(R85) $C_4H_{10} + 1/2O_2 \rightarrow C_4H_8 + H_2O$	$VO_x/Al_2O_3$ , <sup>470</sup> $VO_x/Mg-Al$ spinel, <sup>470</sup>
	1,3-Butadiene	(R86) $C_4H_{10} + O_2 \rightarrow C_4H_6 + 2H_2O$	$VO_x/Mg-Al$ hydrotalcite, <sup>470</sup> $VMgO$ , <sup>470</sup>
	Acetic acid	(R87) $C_4H_{10} + 5/2O_2 \rightarrow 2CH_3COOH + H_2O$	$Mg_3(VO_4)_2-MgO$ , <sup>471</sup> $Mg_2V_2O_7$ , <sup>471</sup> $(VO)_2P_2O_7$ <sup>471</sup>
1-Butene	1,3-Butadiene	(R88) $C_4H_8 + 1/2O_2 \rightarrow C_4H_6 + H_2O$	$V_2O_5/TiO_2$ <sup>472</sup>
	Glutaraldehyde	(R89) $C_5H_8 + O_2 \rightarrow C_5H_6O_2$	$VO_x/Al_2O_3$ , <sup>470</sup> $VO_x/Mg-Al$ spinel, <sup>470</sup>
Ethyl lactate	Ethyl pyruvate	(R90) $C_5H_{10}O_3 + 1/2O_2 \rightarrow C_5H_8O_3 + H_2O$	$VO_x/Mg-Al$ hydrotalcite, <sup>470</sup> $VMgO$ , <sup>470</sup>
Cyclohexane	Cyclohexanone	(R91) $C_6H_{12} + 1/2O_2 \rightarrow C_6H_{10}O + H_2O$	$USb_3O_{10}$ , <sup>473</sup> $USbO_5$ , <sup>473</sup> $MO_3/SiO_2$
	Cyclohexanol	(R92) $C_6H_{12} + 1/2O_2 \rightarrow C_6H_{12}O$	(M = Cr, Mo or W) <sup>474</sup>
Glucose	Lactic acid	(R93) $C_6H_{12}O_6 + O_2 \rightarrow C_3H_6O_3 + 3CO_2 + 3H_2O$	$WO_3/SiO_2$ <sup>475</sup>
5-Hydroxymethyl-furfural	2,5-Furandicarboxylic acid	(R94) $C_6H_6O_3 + 3/2O_2 \rightarrow C_6H_4O_5 + H_2O$	Mesoporous $VO_x/TiO_2$ <sup>476</sup>
Benzene	Maleic acid	(R95) $C_6H_6 + 9/2O_2 \rightarrow C_4H_4O_4 + 2CO_2 + 2H_2O$	$Mn_{0.5}Ce_{0.5}O_x$ <sup>477</sup>
Toluene	Benzaldehyde	(R96) $C_7H_8 + O_2 \rightarrow C_7H_6O + H_2O$	$LaCoO_3$ <sup>478</sup>
	Benzoic acid	(R97) $C_7H_8 + 3/2O_2 \rightarrow C_7H_6O_2 + H_2O$	Co–Mn mixed oxide <sup>479</sup>
Benzyl alcohol	Benzaldehyde	(R98) $C_7H_8O + 1/2O_2 \rightarrow C_7H_6O + H_2O$	$V_2O_5-MoO_3$ <sup>480</sup>
<i>p</i> -Cresol	<i>p</i> -Hydroxybenzyl alcohol	(R99) $C_7H_8O + 1/2O_2 \rightarrow C_7H_8O_2$	$V_2O_5/TiO_2$ , <sup>481</sup> $V_2O_5-MoO_3/TiO_2$ <sup>481</sup>
Ethylbenzene	Styrene	(R100) $C_8H_{10} + 1/2O_2 \rightarrow C_8H_8 + H_2O$	$V_2O_5/TiO_2$ , <sup>481</sup> $V_2O_5-MoO_3/TiO_2$ <sup>481</sup>
<i>o</i> -Xylene	Phthalic anhydride	(R101) $C_8H_{10} + 3O_2 \rightarrow C_8H_4O_3 + 3H_2O$	$Au/CeO_2$ <sup>482</sup>
Naphthalene	Phthalic anhydride	(R102) $C_{10}H_8 + 9/2O_2 \rightarrow C_8H_4O_3 + 2CO_2 + 2H_2O$	MWCNT– $Mn_3O_4$ composite <sup>483</sup>
			$Mn_xTi_{1-x}O_2$ ( $x = 0-0.15$ ) <sup>484</sup>
			$V_2O_5$ <sup>485</sup>
			$V_2O_5$ -Based catalysts <sup>486</sup>





Table 3 (continued)

Feedstock	Products	Reaction	Heterogeneous catalysts
Amines	Imines	(R103) $R/R'-CH_2-NH_2 + O_2 \rightarrow R-CH = NH-CH_2-R/R' + H_2O$ (R = aryl, R' = alkyl)	Meso-Cs/MnO <sub>x</sub> <sup>487</sup>
Methyl bromide	Bromine	(R104) $2CH_3Br + 7/2O_2 \rightarrow 2CO_2 + 3H_2O + Br_2$	Amorphous MnO <sub>x</sub> <sup>488</sup>
1,2-Dichlorobenzene	Chlorine	(R105) $C_6H_4Cl_2 + 7O_2 \rightarrow 6CO_2 + 2H_2O + Cl_2$	Ru/Co <sub>3</sub> O <sub>4</sub> <sup>489</sup>
Formaldehyde	Carbon dioxide	(R106) $CH_2O + O_2 \rightarrow CO_2 + H_2O$	Au/α-MnO <sub>2</sub> <sup>490</sup>
Benzene	Carbon dioxide	(R107) $C_6H_6 + 15/2O_2 \rightarrow 6CO_2 + 3H_2O$	MCo <sub>2</sub> O <sub>4</sub> (M = Cu, Ni or Mn) <sup>491</sup>
Toluene	Carbon dioxide	(R108) $C_7H_8 + 9O_2 \rightarrow 7CO_2 + 4H_2O$	LaMnO <sub>3</sub> , <sup>492</sup> La <sub>0.9</sub> Sr <sub>0.1</sub> MnO <sub>3</sub> , <sup>492</sup> La <sub>0.9</sub> Sr <sub>0.1</sub> Mn <sub>0.9</sub> Fe <sub>0.1</sub> O <sub>3</sub> , <sup>492</sup> Ag, Pt or Pd doped CoAl layered double hydroxide <sup>493</sup>
Ammonia	Nitrogen	(R109) $2NH_3 + 3/2O_2 \rightarrow N_2 + 3H_2O$	Fe <sub>2</sub> O <sub>3</sub> , <sup>494</sup> Cr <sub>2</sub> O <sub>3</sub> , <sup>494</sup> CeO <sub>2</sub> <sup>494</sup>
	Nitric oxide	(R110) $NH_3 + 5/4O_2 \rightarrow NO + 3/2H_2O$	Fe <sub>2</sub> O <sub>3</sub> , <sup>494</sup> Cr <sub>2</sub> O <sub>3</sub> , <sup>494</sup> CeO <sub>2</sub> <sup>494</sup>
Hydrogen sulfide	Sulfur	(R111) $H_2S + 1/2O_2 \rightarrow 1/nS_n + H_2O$	La <sub>1-x</sub> Ce <sub>x</sub> FeO <sub>3</sub> (x = 0, 0.2, 0.4, 0.6 and 0.8), <sup>495</sup> Cu-V and Cu-V-Mo mixed oxides <sup>496</sup>

carriers, also offer exciting potential for chemical syntheses. Aside from these, significant progress still needs to be made towards in-depth understanding of redox reaction mechanisms, catalyst design and optimizations, reactor and process design and operations, as well as process scale up for the existing CLBC approaches. To summarize, we believe that CLBC represents a largely uncharted and extraordinarily exciting area of research and potential applications.

## Conflicts of interest

Authors have no conflict of interest to declare.

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