



Cite this: *Energy Adv.*, 2023, 2, 1476

A novel technological blue hydrogen production process: industrial sorption enhanced autothermal membrane (ISEAM)

Chidozie Eluwah, ^{ab} Paul S. Fennell,^a Christopher J. Tighe^a and Ahmed Al Dawood ^b

A novel technological industrial blue hydrogen production process – the Industrial Sorption Enhanced Autothermal Membrane (ISEAM) process, with the potential to produce constant fuel cell grade hydrogen with a purity of 99.99%, regardless of upstream process upsets, has been modelled using an Aspen Plus simulator and MATLAB (including both thermodynamics and kinetics analysis). The process exhibits a very high hydrogen yield (99%), and methane conversion (99.9%), with a low carbon monoxide footprint (at ppm levels). The results were validated by comparing against experimental data published in the literature. Parametric evaluations were later conducted to identify the optimal operating conditions for the developed blue hydrogen ISEAM process. The required reforming heat is provided by the exothermic carbonation reaction of a sorbent, while chemical looping of the oxygen carrier (metal oxides) provides the regeneration heat required for the saturated sorbent, in a novel multi-tubular packed shell and tube reactor. Pinch analysis shows that the process is auto thermal (so it does not need any external heating utility) and can achieve an extremely high 97.5% thermal and hydrogen production efficiency. The ISEAM process was benchmarked against an industrial steam methane reforming (SMR) plant and the result shows $\geq 32\%$ improvements in most of the technical parameters that were evaluated. Economic evaluation shows a levelized cost of hydrogen (LCOH) of \$2.6 per kg-H₂ for the baseline SMR plant compared with \$1.3 per kg-H₂ for the ISEAM process (a 50% cost reduction). The cost of CO₂ removal (CCR) was calculated as \$180 per tonneCO₂ for the baseline SMR process compared with \$33.2 per tonneCO₂ (81.6% cost reduction) for the novel process. The novel ISEAM process utilizes mature and existing industry technologies such as desulphurization, pre-reforming, adsorption, membranes, waste heat boilers, and pressure swing adsorption. Because of this, scale-up is easier and some of the challenges associated with the SMR process and integrated sorption enhanced membrane reforming (SEMR) processes are addressed. These include thermodynamic constraints, a high energy penalty, overall process integration, optimization, membrane contamination, carbon deposition and unsteady state operation.

Received 23rd February 2023,
Accepted 21st July 2023

DOI: 10.1039/d3ya00090g

rsc.li/energy-advances

1. Introduction

The burning of fossil fuels and emission of associated greenhouse gases (GHG) into the atmosphere poses a serious threat to the global environment, and contributes to climate change. To meet the United Nations Climate Change target of limiting global warming to 1.5 °C above pre-industrial levels, a drastic reduction in CO₂ emissions and significant changes in the energy sector are needed. According to the US Energy

Information Administration,¹ as of January 1, 2020, there are approximately 206 trillion cubic metres of total world reserves of natural gas. Hydrogen has a vital part to play in the global future energy mix^{2,3} because of its zero-carbon combustion and its high energy per unit mass (though admittedly, low energy per unit volume at STP). It is estimated that 50% of hydrogen produced worldwide is produced using Steam Methane Reforming (SMR).⁴ This process is inefficient and thermodynamically limited; it is also a large emitter of CO₂, estimated at almost 2% of the global CO₂ emissions in 2019.⁵ Hence, the deployment of an environmentally friendly and economically attractive process⁶ with a low energy-penalty is of huge interest to researchers. Various research studies have been carried out to address the limitations of steam methane reforming (SMR)^{7,8}

^a Department of Chemical Engineering, South Kensington Campus, Imperial College London, London SW7 2AZ, UK

^b Unconventional Resource Production Department, Saudi Arabian Oil Company, Saudi Arabia



which include Sorption Enhanced Reforming (SER),^{9–29} Membrane Reforming (MR),^{17,25,30–33} integrated Sorption Enhanced Membrane Reforming (SEMR)^{34,35} and Gas Switching Reforming (GSR). Abanades *et al.*^{24,36–38} proposed the use of Ca/Cu looping in a single reactor through a three-step process; (a) sorption enhanced reforming with simultaneous carbonation of CaO; (b) oxidation of Cu to CuO with air; (c) calcination of CaCO_3 during the reduction of CuO with a fuel gas.³⁹ The exothermic carbonation of CaO and the exothermic oxidation of Cu to CuO supply the required reforming and calcination heat respectively. *In situ* removal of CO_2 during the reforming step increased the methane conversion. One of the main practical challenges of Ca/Cu looping which this current paper addresses in the SER unit of the novel process, is the elimination of the potential venting of CO_2 to the atmosphere as a result of decomposition of CaCO_3 during the oxidation step. Shahid *et al.*⁴⁰ investigated the sorption-enhanced reforming (SER) performance in a packed bed reactor for three CO_2 sorbents such as calcium oxide (CaO), lithium zirconate (LZC) and hydrotalcite (HTC). They concluded that CaO-based sorbents exhibited optimum conditions of 627 °C and 300 kPa with methane conversion of 82% and hydrogen purity of 85%. LZC and HTC were suitable under optimal conditions of 500 kPa and 500 °C with a methane conversion range of 91.2–94.1% and hydrogen purity of 55.1–77.8%, which exceeds the performance of the SMR process under similar operating conditions. Diglio *et al.*⁴¹ investigated SER with 1D numerical modelling in a fixed bed reactor network, packed with the Ni-catalyst/Ca-sorbent operated under cyclical carbonation/reforming and calcination conditions using a pressure swing at 700 °C. During the carbonation/reforming mode, the reactor's feed was a mixture of methane and steam at a high pressure of 3500 kPa. To shift the equilibrium to calcination, the reactors were fed with a mixture of steam and CO_2 at a reduced reactor pressure of 100 kPa. Six stages of cyclic operation of the reactor were required: (1) carbonation (CS); (2) purging (PS) for hydrogen removal; (3) depressurisation (D) to obtain a vacuum condition in the bed; (4) calcination (CAS); (5) purging for removal of steam and CO_2 and (6) pressurisation (PR) to raise the pressure to the carbonation/reforming pressure of 3500 kPa. The results showed that at least 8 reactors in parallel were required to produce a hydrogen purity of 92% with an average hydrogen yield of 2.9 molH₂ per molCH₄. Nazir *et al.*⁴² developed a gas switching reforming (GSR) process where a cluster of several standalone reactor cycles through three steps: oxygen carrier reduction by PSA off-gas, steam methane reforming, and oxidation with air. The GSR hydrogen process was claimed to have 3% higher equivalent hydrogen production efficiency and an efficiency penalty of 0.3% for 96% CO_2 capture when compared to SMR. Silva *et al.*⁴³ through dynamic mathematical modelling showed that a methane conversion of 99.85% and a hydrogen yield of 1.626 molH₂ per mol CH₄ (steady state) could be achieved at a temperature of 550–600 °C in a fixed bed membrane reactor because of *in situ* removal of hydrogen relative to a methane conversion of 88.87% and a hydrogen yield of 1.445 molH₂

per mol CH₄ (steady state) at 725–950 °C for a conventional SMR reactor. Kim *et al.*⁴⁴ developed an industrial high permeable Pd-alloy membrane which was prepared on tubular-porous stainless steel (PSS) 12.7 mm in diameter and a membrane thickness of 6 μm using electroless plating (ELP) of Pd and Ru. Hydrogen permeance after a long-term reforming test was 0.00346 Mol m^{–2} Pa^{0.5} which is quite high compared with other reports. Ye G. *et al.*⁴⁵ and Shafiee *et al.*⁴⁶ modelled an MR using a sequential model with sub-membrane reactors and sub-separators. Parametric investigations showed that *in situ* hydrogen removal breaks the thermodynamic constraints and improved hydrogen yield and methane conversion, especially at a high permeation capacity. Lee *et al.*⁴ developed an integrated SEMR process with membrane and sorption enhancement in a single reactor. Techno-economic simulation showed that simultaneous *in situ* removal of CO_2 and hydrogen from the reaction zones produces the highest methane conversion and hydrogen yield compared with Membrane Reactor (MR) only, Sorption Enhanced Reactor (SER) only, and an SMR process. Ji *et al.*⁴⁷ used Computational Fluid Dynamics (CFD) to prove that SEMR not only decreased the CO_2 emission and CO poisoning of the catalyst and membrane, but also increased the hydrogen yield, methane conversion and reaction kinetics for both the reforming and water gas shift reactions. The CO mol fraction decreased by 1 order of magnitude in the SEMR compared to the traditional membrane reactor, which minimized the possibility of hydrogen permeation decay.

Dou *et al.*⁴⁸ proposed a sorption-enhanced chemical looping steam reforming of glycerol with CO_2 *in situ* capture and utilization through a dry methane reforming process (SE-CLSR + MDR). The process was operated in three stages. The first stage is a typical sorption-enhanced chemical steam reforming process at 650 °C, utilizing $\text{NiO}/\text{NiAl}_2\text{O}_4$ as catalysts, calcined dolomite as the sorbent and glycerol as the feedstock. The second stage process was the simultaneous calcination of CaCO_3 at 850 °C and dry methane reforming by the use of methane feedstock using the reduced $\text{Ni}/\text{NiAl}_2\text{O}_4$ as catalysts. The last stage included the oxidation stage where the deposited carbon on the catalyst was oxidised using pure oxygen and the reduced nickel catalyst is re-oxidised to nickel oxide. The results demonstrated that hydrogen of above 92.5% could be generated through a one-step process and CO_2 decreased to zero in the pre- CO_2 breakthrough periods improving the economics and lower impact on the environment. The proposed process (SE-CLSR + MDR) has some similarities with the novel ISEAM process such as use of calcined dolomite for *in situ* CO_2 removal, chemical looping technology and the use of multiple reactors simultaneously in order to operate the process in a steady state. However, the ISEAM process differs in the main feedstock (methane), process integration, higher feedstock conversion and yield, application of chemical looping combustion which is only used to generate the required calcination heat through an in-direct heat transfer with the main process, the use of a hydrogen-permselective palladium membrane reactor to drive the overall conversion to 99.9%, the use of air



in a novel multi-tubular reactor with no risk of diluting the recovered CO₂ during the calcination stage as against the use of pure oxygen in the Dou *et al.* process which requires energy intensive and a costly air separation unit.

The goal of this paper is to propose a novel industrial sorption enhanced autothermal membrane (ISEAM) process as an alternative to the leading industrial steam methane reforming process. The novel process addresses the limitations of the SMR process, such as thermodynamic equilibrium constraints, catalyst deactivation and high energy penalty. In addition, it addresses problems associated with the sorption enhanced membrane reforming (SEMR) process, such as membrane contamination, carbon deposition, unsteady state operation and overall process integration.

2. Methodology

2.1 Baseline SMR process description

Fig. 1 shows a block flow diagram of a baseline SMR plant with an amine unit integrated to remove the CO₂. Natural gas as per Table 1 and steam are converted to H₂ and CO in a steam reforming process under severe conditions of temperature of 800–1000 °C and pressure of 2000–3500 kPa in the presence of a nickel alumina catalyst.

The steam reforming stage is followed by two water-gas-shift (WGS) reaction steps to maximize the conversion of CO to H₂ and by a final pressure swing adsorption (PSA) unit to produce hydrogen with a purity of 99.9%. An amine unit is added downstream of the water gas shift (WGS) reactor and reaction furnace (RF) to capture the CO₂ produced from the process and the flue gas.

Table 1 Baseline plant data⁴⁹

Components	Mole (%)
C1	77.42
C2	13.27
C3	5.76
CO ₂	2.29
N ₂	0.63
n-C4	0.36
i-C4	0.23
n-C5	0.02
i-C5	0.02
Feed gas, kmol h ⁻¹	287.6
S/C	2.9

2.2 Novel process description

As shown in Fig. 2 and 3, natural gas with feed compositions as per Table 1 is compressed to 3600 kPa and 93 °C and heated through a shell and tube heat exchanger to 380 °C before being fed into a feed gas pre-treatment unit where organic sulfur is removed and heavy hydrocarbons are converted into a mixture of methane, carbon dioxide, carbon monoxide and hydrogen over a nickel oxide catalyst.⁵⁰ Desulphurization of the feed gas is required to prevent poisoning of the reformer catalyst. The pre-conditioned natural gas feedstock is heated through a series of heat exchangers using the hot effluent retentate and permeate gas from the downstream Pd-alloy membrane catalytic reactor. This is mixed with HP steam produced from the waste heat boiler (WHB). The combined natural gas and HP steam is superheated to 500 °C and routed to the multi-tubular adiabatic packed bed Carbonator-SMR Catalytic Reactor. The reactor is a shell and tube, with packed solid sorbent such as CaO and Ni/Al₂O₃ catalyst in the tube side and packed oxygen carrier (metal oxides) such as CuO/Al₂O₃ in the shell side. The

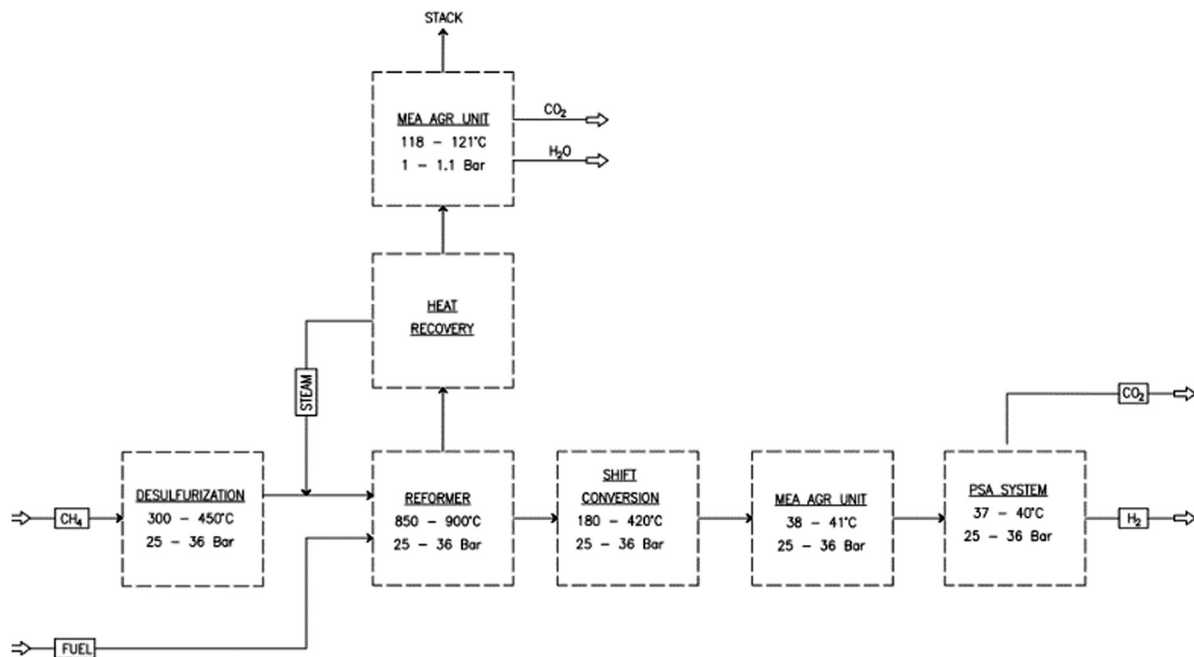


Fig. 1 Block flow diagram of the steam methane reforming (SMR) process

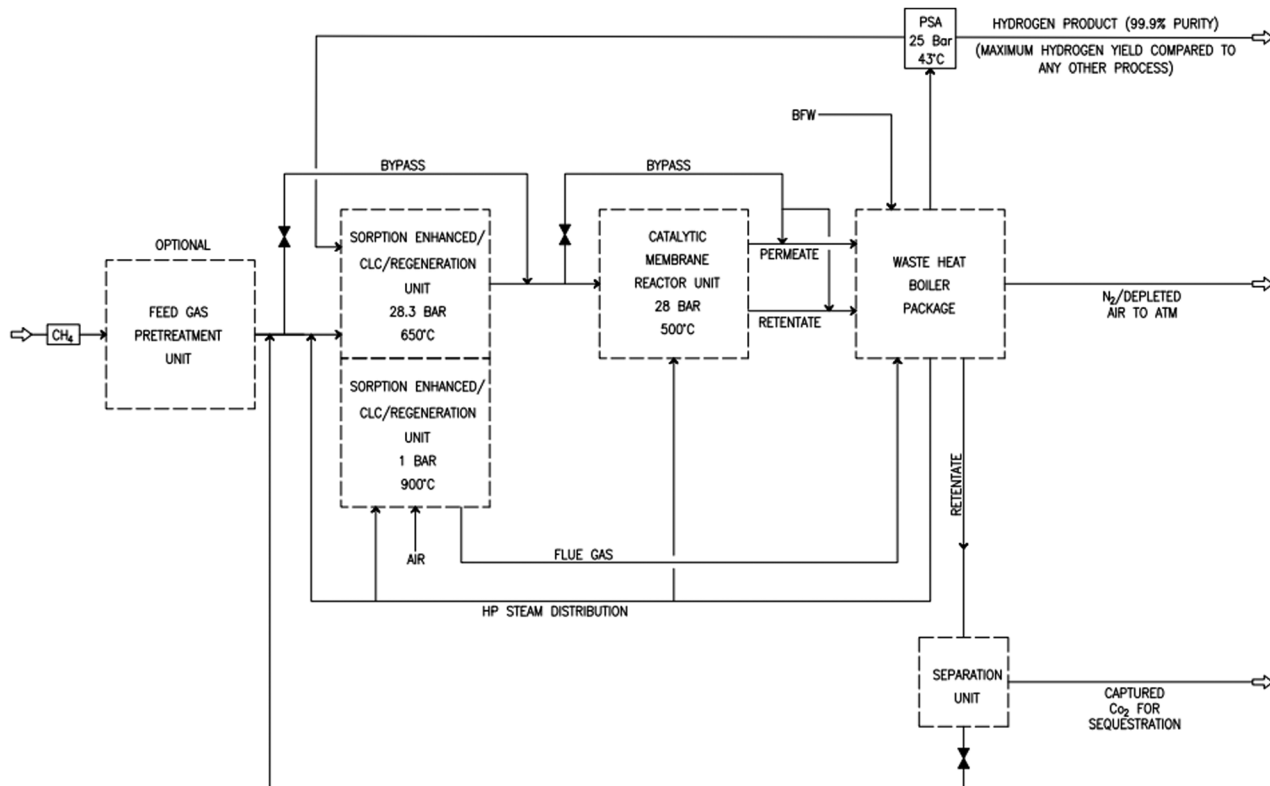
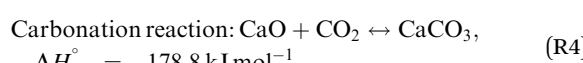
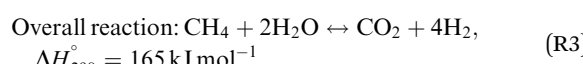
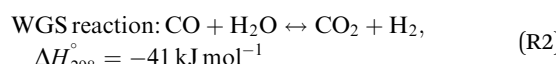
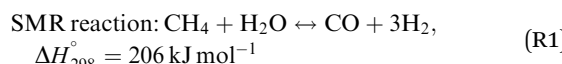


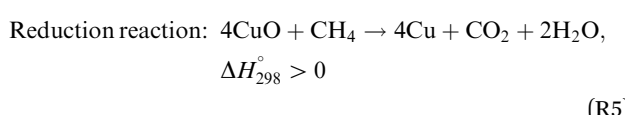
Fig. 2 Novel industrial sorption enhanced auto-thermal membrane (ISEAM) process.

Carbonator-SMR Reactors are $3 \times 100\%$ regenerative packed bed reactors, with one in the adsorption-reforming mode, the second in the regeneration-oxidation mode and the third in the cooling/stand-by mode. The reactions in the tube side of the Carbonator SMR Reactor are auto thermal as the exothermic carbonation reaction produces the required heat for the endothermic reforming reactions.

Tube side reactions:



Shell side:

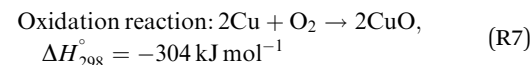


The main reactions that occur in the Carbonator-SMR catalytic Reactor during the regeneration-oxidation mode are:

Tube side:



Shell side:



In situ removal of CO₂ in the Carbonator-SMR Reactor shifts the reforming and water gas shift reactions to the forward direction as per Le Chatellier's principle, leading to an increased hydrogen yield and methane conversion. The use of a sorption enhanced reforming unit has the potential to introduce unsteady state transient operation. Fig. 4 shows a typical behaviour of the Carbonator-SMR Reactor. Three typical periods of interest are identified: period 1 between $0 < t < 1000$ s typical, transient period between $1000 < t < 2000$ s, and period 2 $t > 2000$ s. Period 1 ($0 < t < 1000$ s) is the pre-breakthrough stage, which is characterized by fast kinetics as the active surface area of the sorbent (CaO) is available leading to higher hydrogen production. The transient period is the breakthrough stage ($1000 > t < 2000$ s). This stage is characterized by slow kinetics as the carbonation reaction is controlled by the diffusion through the solid product (CaCO₃) layer. The CaO conversion rate is reduced as more product (CaCO₃) is formed and deposited on the active surface of the CaO. This leads to a decline in the product hydrogen. The last stage is the post-breakthrough ($t > 2000$ s); during this stage, the CaO sorbent is saturated and hence there is a total loss of sorption, and the

article is licensed under a Creative Commons Attribution-NonCommercial 3.0 Unported Licence.
Published on 1 August 2023. Downloaded on 21/10/2026 2:58:33 PM.

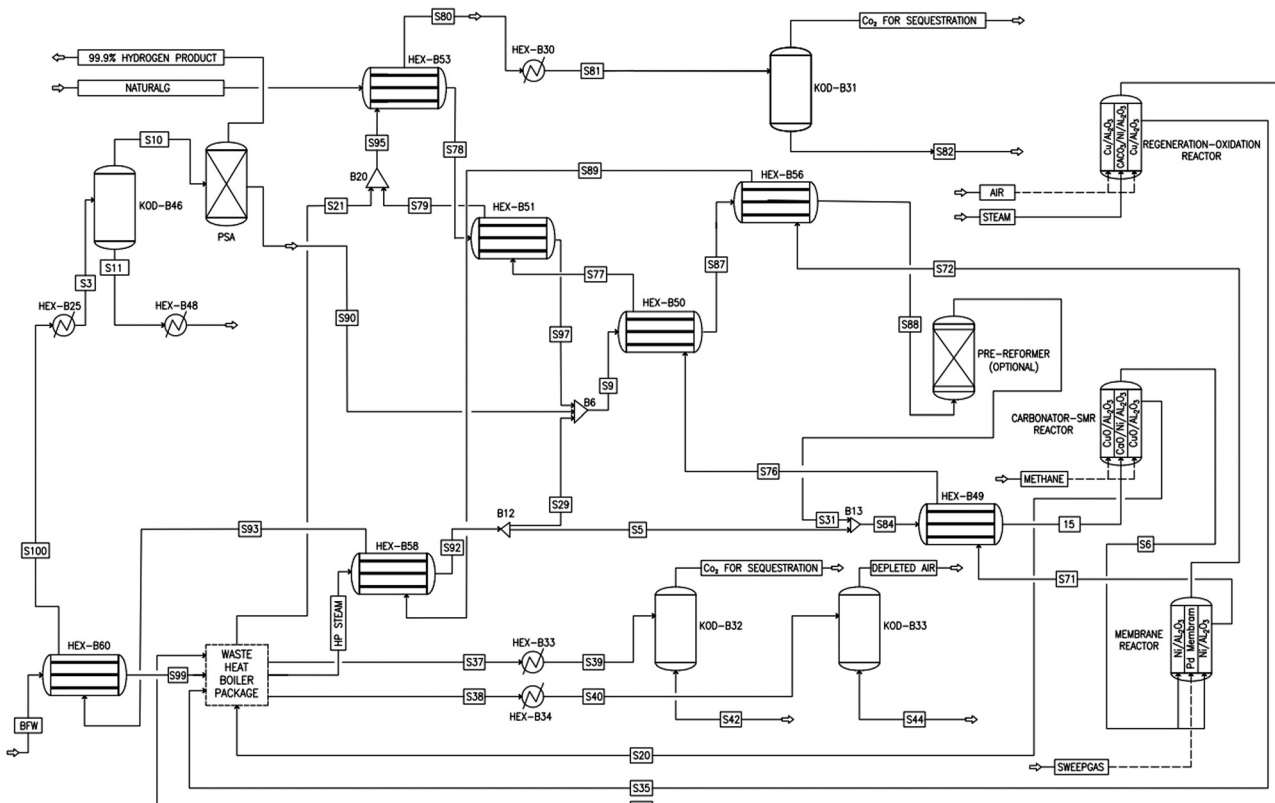


Fig. 3 Process flow diagram of a novel industrial sorption enhanced auto-thermal membrane (ISEAM) process.

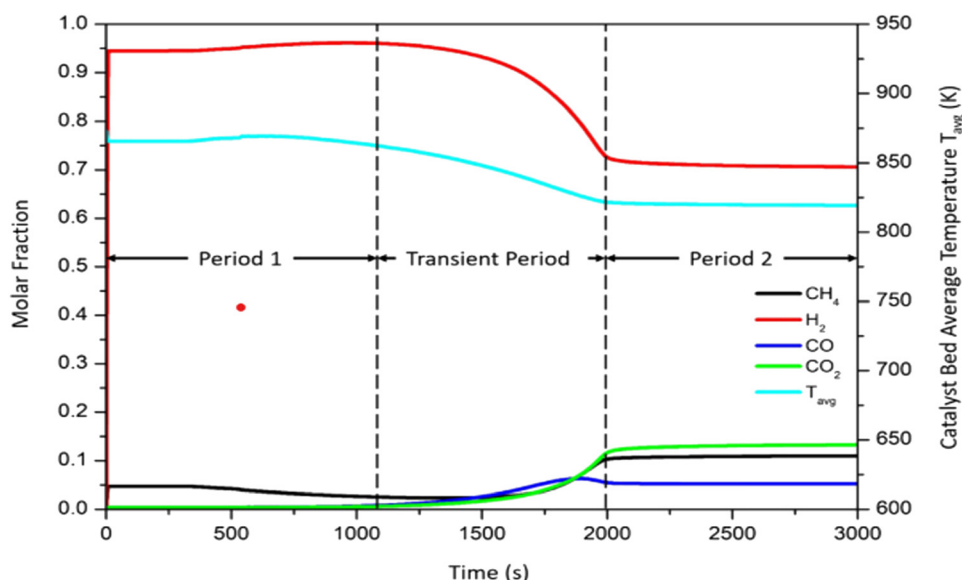


Fig. 4 Typical product plot for 100% sorption enhanced reforming process.⁵¹

reactions return to those of conventional steam methane reforming.

To run the $3 \times 100\%$ Carbonator-SMR Reactor in the steady state mode, switching between adsorption and calcination is carried out within period 1. This can be done automatically

based on time or started by the reactor outlet gas analysers depending on the CO₂ concentration in the product gas. The product gas mixture from the tube side of the Carbonator-SMR Reactor leaves at a temperature of 580–650 °C before being routed to the Pd-Alloy catalytic Membrane Reactor, packed with

the Ni/Al₂O₃ catalyst, where further reforming and water gas shift reactions occur. The reforming heat in the Pd-alloy membrane Reactor is provided by the sensible heat of the gas from the upstream Carbonator-SMR Reactor. Steam is used as a sweeping gas to enhance the permeation of hydrogen through the Pd-Alloy Membrane Reactor. Like the Carbonator-SMR Reactor, in-situ removal of the product hydrogen from the reaction zones of the Membrane Reactor results in a further increase of hydrogen yield and methane conversion in accordance with reactions (R1)–(R3). The permeated gas from the Membrane Reactor contains mainly hydrogen and steam, which is used to pre-heat the feed gas and boiler feed water. To achieve a continuous high quality fuel cell hydrogen purity grade of 99.99% regardless of the transient behavior of the process, the cooled permeating gas is routed through a Pressure Swing Absorber (PSA) for final purification and polishing. The retentate gas from the Pd-alloy membrane contains mainly CO₂ and steam, which is separated by cooling and the use of a knock-out drum. The heat required for calcination is provided by the exothermic heat of the chemical looping combustion of Cu/CuO or any other oxygen carriers such as NiO and FeO in the presence of an Al₂O₃ catalyst. The sorption enhanced reforming (SER) unit of the ISEAM process is different from the conventional Ca/Cu looping proposed by Abanades *et al.*³⁸ as the sorbent in the novel process is charged into a novel multi-tubular section of the Carbonator-SMR Reactor while the oxygen carrier (metal oxide) is loaded in the shell side of the reactor, providing the required calcination heat through indirect heating; thus, eliminating the problem of potential venting of CO₂ to the atmosphere as a result of decomposition of CaCO₃ during the oxidation step associated with conventional Ca/Cu looping. The process uses mature and existing industry technologies such as a desulphurization unit, pre-reforming unit, adsorption, membrane unit, waste heat boiler and PSA unit.

2.3 ISEAM process heat integration

Pinch analysis is an energy optimisation technique where cold and hot streams are matched for the purpose of optimum utilization of the process heat energy, and enables the determination of the minimum utility heating and cooling duties required by the process. This makes it possible to design an optimal heat exchanger network, reducing both capital and operation costs. Pinch analysis was conducted using an Aspen Energy Analyser for the ISEAM process. A minimum temperature approach of 10 °C was used. A required minimum process temperature of 37 °C is dictated by the PSA unit and the maximum process temperature of 900 °C is dictated by the regeneration unit of the Carbonator SMR Reactor. The heat needed for regeneration (calcination) is provided by the exothermic chemical looping combustion reaction as per reaction (R7). The external cooling water supply temperature is taken as 20 °C. Significant opportunities for heat integration were identified as shown in Fig. 6 and briefly outlined below for the main streams:

(1) Hot effluent gas (mostly depleted air) at a temperature of 960 °C from the shell side of the regenerator-oxidation reactor (air reactor) is used in the waste heat boiler to produce process steam from boiler feed water.

(2) Hot effluent gas (mostly CO₂) at a temperature of 928 °C from the tube side of the regenerator reactor (fuel reactor) is used in the waste heat boiler to produce process steam from boiler feed water.

(3) Hot permeate product gas at a temperature of 500 °C from the Pd-alloy membrane reactor (stream S72) is matched with cold pre-reforming feed gas (stream S87).

(4) Hot retentate product gas (stream S71) at a temperature of 500 °C from the Pd-alloy membrane is used to preheat the feed gas (stream S84) to the SMR-Carbonator Reactor. Fig. 5 shows the composite curve and Fig. 6 shows the heat exchangers grid diagram.

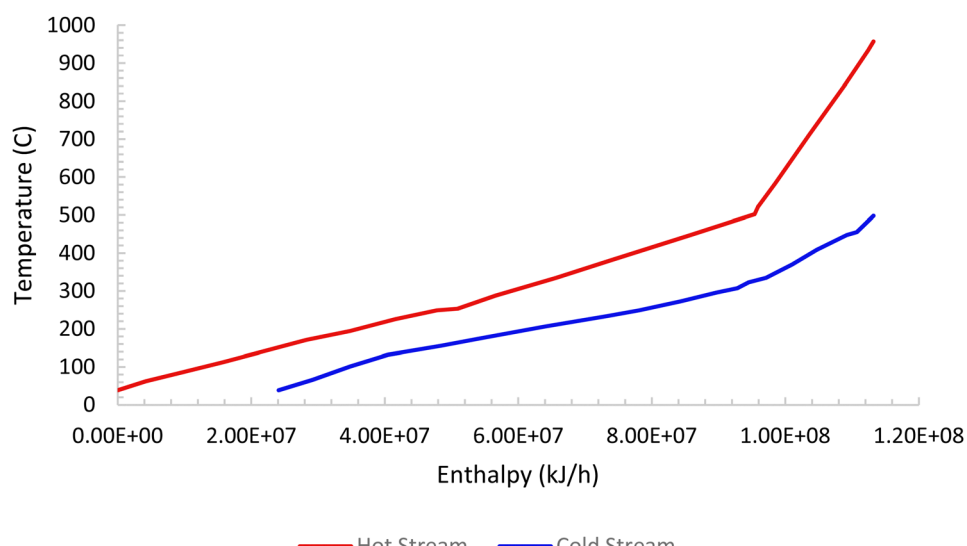


Fig. 5 ISEAM process composite curve.



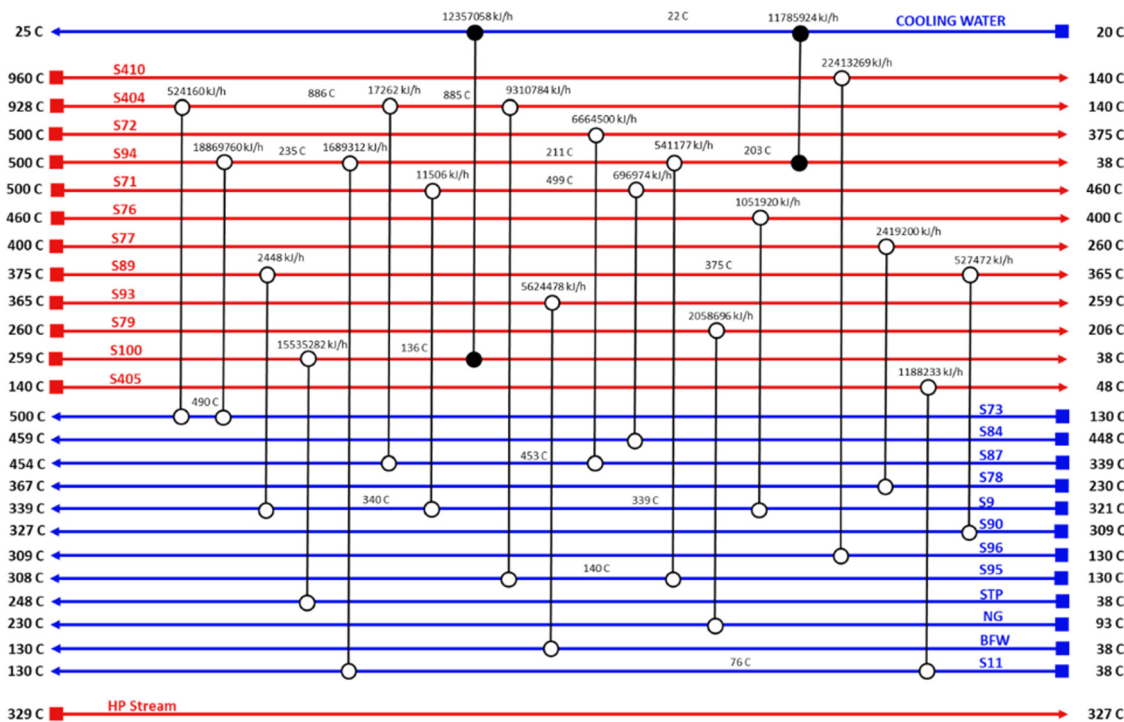


Fig. 6 ISEAM process heat exchanger grid diagram.

(5) As shown in the heat exchanger grid diagram (Fig. 6), there is no external added heating utility required, making the process auto thermal.

2.4 Process modelling and simulation

A steady state thermodynamic equilibrium and kinetic model was developed using integrated Aspen Plus and MATLAB software and used to model and evaluate the baseline SMR plant and the novel process based on a 1-D heterogenous reactor.

model. Data in Table 2 was used for the novel ISEAM process modelling. Steam methane reforming rate kinetic models (Table 11 in Appendix B) developed by Xu *et al.*⁵² in the presence of a Ni/Al₂O₃ catalyst based on the Langmuir–Hinshelwood–Hougen Watson (LHHW) mechanism was adopted and used for the kinetic modelling. Carbonation kinetic rate and data was adopted from a study by Rodriguez *et al.*⁵³ The hydrogen permeation through the membrane was modelled using Sievert's law (eqn (27) in Appendix B), and the membrane

Table 2 System and operating specifications for the novel ISEAM process

Parameter	Value	Units
Membrane reactor		
Number of reactors	$2 \times 50\%$	—
Reactor pressure	2800	kPa
Reactor inlet temperature	580–650	°C
Permeate inlet pressure	2500	kPa
Membrane tube diameter	0.09	M
Membrane thickness	6	μm
Tube length	23.5	M
Number of tubes	80	—
Pressure difference between permeate and retentate	250	kPa
Membrane permeance ⁴⁴	0.00346	Mol m ⁻² s Pa ^{0.5}
Carbonator SMR reactor		
Number of reactors	$3 \times 100\%$	—
Reactor pressure	2800	kPa
Reactor inlet temperature	500	°C
Tube diameter	0.1143	m
Tube length	7	m
Number of tubes	40	
Reactor temperature	500	°C

Table 3 Parameters and assumptions for economic analysis (CAPEX)

Equipment	Number required	Bare equipment cost (BEC), \$1000	Ref.
SMR reactor	1	57 841	60
Sulfur polisher (ZnO)	1	243	60
Primary air compressor	2(1)	861	60
Water gas shift reactor	4	12 918	60
MDEA CO ₂ unit (95% CO ₂ removal)	2	95 985	60
MEA CO ₂ unit (90% CO ₂) removal	2	68 176	60
PSA unit	2	38 047	60
Boiler package	1	7306	60
Carbonator (k€2017)	1	$474. \text{ Transferred thermal power (MW}_{\text{th}}\text{)} + 8360. \left(\frac{\text{Internal diameter (m)}}{4.7} \right)^{0.6}$	59
Calciner (k€2017)	1	$415. (\text{Fuel thermal input (MW}_{\text{th}}\text{)})^{0.65}$	59
Membrane reactor	1	\$4203.75/surface area (m ²)	4

parameters and permeance developed by Kim *et al.*⁴⁴ were used because of the high permeance value and suitability for industrial application.

2.5 Economic evaluation

Economic evaluation of the baseline SMR process and the novel ISEAM process (base and optimised) were conducted following the method proposed by the Global CCS Institute^{54,55} using key economic indicators such as the cost of CO₂ avoided (CCA), cost of CO₂ removal (CCR) and levelized cost of hydrogen (LCOH).⁵⁶ These costs were calculated using eqn (1)–(5).^{5,42} The fixed charge factor (FCF) converts the total capital requirement into uniform annual amounts at a discount rate over the lifetime of the plant. Total capital requirement (TCR), commonly referred to as CAPEX, is calculated using the installed cost for main equipment and the assumptions stated in Tables 3 and 4. Balance of Plant (BOP) cost include the cooling system, fuel, electricity, storage, make-up water, sanitary system, water discharge and solid wastes, *etc.* Bare equipment costs were obtained from various works^{5,42,57–61} and adjusted to the year 2022 using eqn (5),⁵ and the IHS global capital cost escalation index factor was applied. Assumptions used for the fixed operating and maintenance cost (FOM) and variable operating

and maintenance cost (VOM) are summarised in Table 5.

$$\text{LCOH} = \frac{(\text{TCR})(\text{FCF}) + \text{FOM}}{(\text{M}_{\text{H}_2})(\text{CF} \times 8760)} + \text{VOM} + (\text{FC})(\text{HR}) \quad (1)$$

$$\text{CCA} = \frac{\text{LCOH}_{\text{with CCS}} - \text{LCOH}_{\text{ref (non CCS)}}}{\text{CO}_2 \text{ Emission (ref non CCS)} - \text{CO}_2 \text{ Emission (with CCS)}} \quad (2)$$

$$\text{CCR} = \frac{\text{LCOH}_{\text{with CCS}} - \text{LCOH}_{\text{ref (non CCS)}}}{\text{CO}_2 \text{ Removed (with CCS)}} \quad (3)$$

$$\text{FCF} = \frac{r(1+r)^r}{(1+r)^r - 1} \quad (4)$$

$$C_A = \left(\frac{C_{I_A}}{C_{I_B}} \right) C_B \frac{(S_A)^X}{S_B} \quad (5)$$

3. Results and discussion

3.1 Model validation

The baseline kinetic model of the SMR process was validated using actual plant data adopted from Salem *et al.*⁴⁹ The

Table 4 Parameters and Assumptions for Economic analysis (CAPEX)

Equipment	Number required	Bare equipment cost (BEC), \$1000	Ref.
Capital cost escalation factor	2007:170 2017:182 2020:205 2022:210		IHS Markit (downstream)
Capacity factor (CF)	0.95		5
Plant design life	25 years		5
Discount rate	12%		5
Engineering, procurement, and construction cost (EPCC)	8% of bare erected cost (BEC)		42
Process contingency	30% of BEC for MDEA unit, MEA unit, carbonator-reactor and membrane reactor; 0% for reference SMR plant		42
Project contingency	10% of (BEC + EPCC + process contingency)		42
Balance of project (BOP)	15% of (BEC + project contingency)		60
Total contingencies	Project contingency + process contingency		42
Total plant cost (TPC)	BEC + EPCC + total contingencies + BOP		42
Owner's cost	20.2% of TPC		42
Total overnight cost (TOC)	TPC + owner's cost		42
Total capital requirement (TCR)	$1.14 \times \text{TOC}$		42



Table 5 Parameters and assumptions for economic analysis (OPEX)

Description	Values	Ref.
Operating labour	\$60000 per person-year	42
Operator per shift	16	60
Total shift per day	2	60
Maintenance, support, and administration	2.5% TOC	42
Property taxes and insurance	2% TOC	42
Fuel cost	\$2.58/MMBTU	4
Variable operating and maintenance cost (VOM)	10% of TOC	

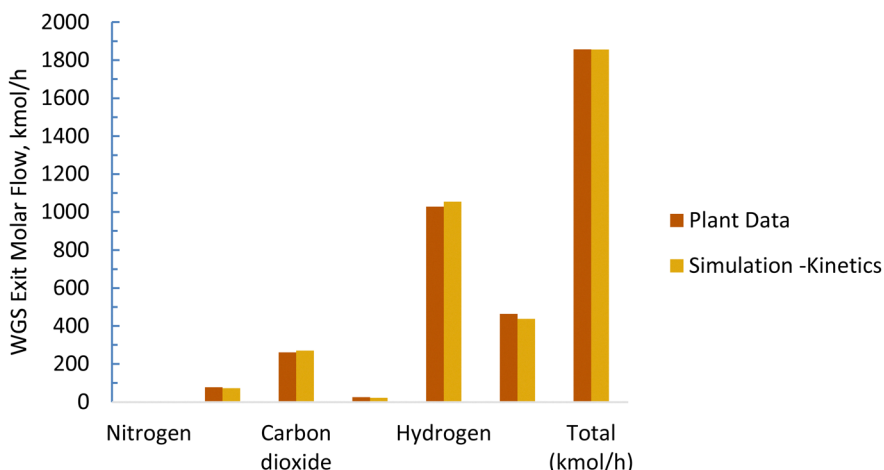


Fig. 7 Baseline WGS model validation.

validation results for the baseline SMR process (Fig. 7 and 8) show that the kinetic model compares well with the actual plant data, with the kinetic model predicting approximately 3% higher methane conversion and hydrogen yield. The sorption enhanced reforming unit model for the ISEAM process was validated using experimental data (Table 9) reported by Lee *et al.*⁶² and the model for the Pd-alloy membrane was confirmed using experimental data (Table 10) reported by Kim *et al.*⁴⁴ The ISEAM kinetic model was validated using the

thermodynamic equilibrium model as shown in Fig. 9 and kinetics model results compare very favourably with the equilibrium results.

3.2 Thermodynamic evaluation of the novel ISEAM process and baseline SMR process

The novel ISEAM process was compared with the baseline industrial SMR process in terms of methane conversion, hydrogen yield, hydrogen purity, hydrogen thermal efficiency and

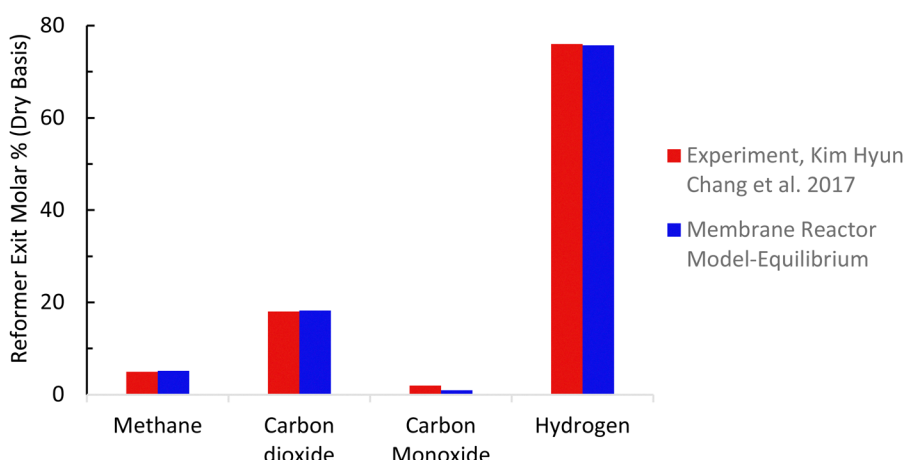


Fig. 8 Membrane reactor model validation.



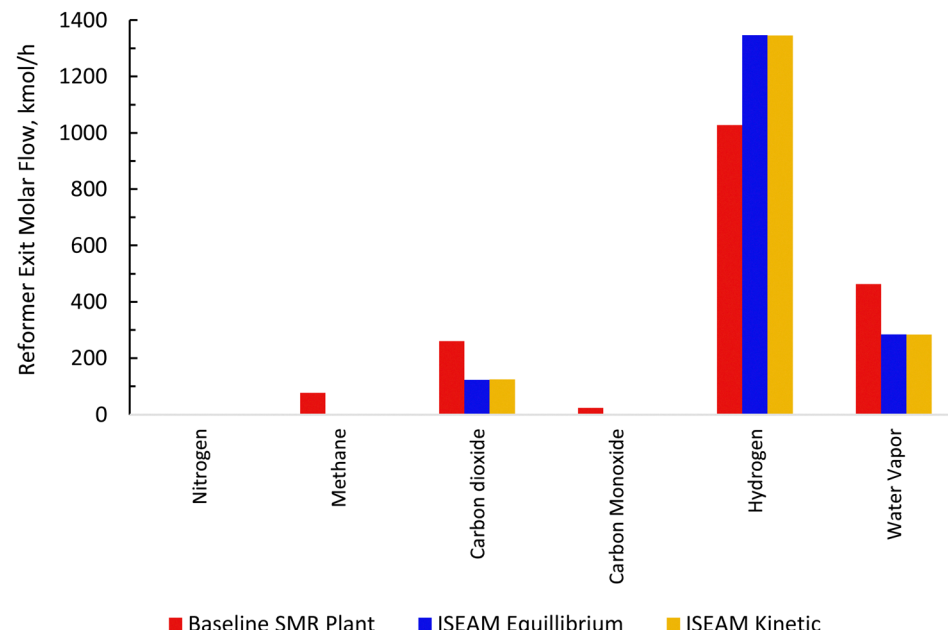


Fig. 9 Comparison between the baseline SMR and novel ISEAM Process plant and novel ISEAM.

process thermal efficiency using eqn (6)–(9),^{40,63,64} where ‘*n*’ stands for relevant molar flowrates. Heating values for 100% natural gas (CH_4) and 100% hydrogen gas are simulated in Aspen Plus with standard conditions of $891.6 \text{ MJ kmol}^{-1}$ and 286 MJ kmol^{-1} respectively. The results presented in Fig. 9, 10 and Table 6 show that the methane conversion and percentage hydrogen yield is higher by 32% for the novel ISEAM process compared with the baseline SMR process as the thermochemical reversible equilibrium constraints that limit the reforming and water gas shift reactions is broken by the in-situ CO_2 and

hydrogen removal. The process thermal efficiency is higher by 41% and the hydrogen production efficiency is higher by 15.8% for the novel process since there is no external heating utility required as the process is fully heat integrated. The process thermal efficiency achieved in the ISEAM process (97.5%) is among the highest compared with typical values of 48–80% reported in the literature for other alternative processes.^{42,57,58} A higher thermal efficiency is achieved using chemical looping combustion (CLC) which provides the regeneration heat, careful heat integration, and CO_2 removal (mainly) by the use of a

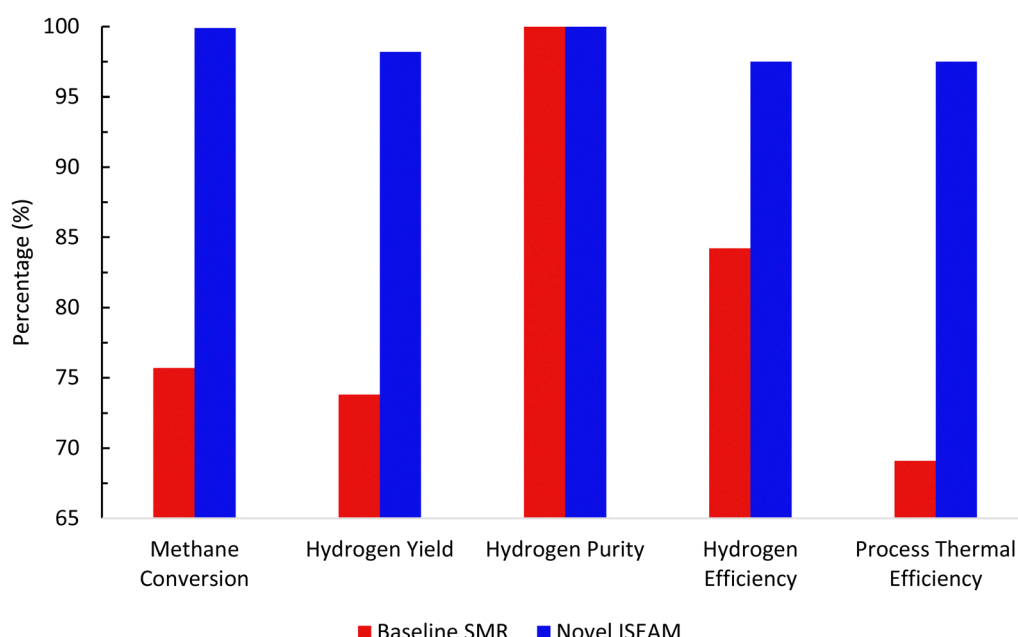


Fig. 10 Comparison between the baseline SMR process and novel ISEAM process.

Table 6 Summary of the results comparing baseline plant data and the novel ISEAM process

Description	Operating temperature, °C	Net heating utility, MJ s ⁻¹	Total thermal energy input, MJ s ⁻¹	Total thermal energy output, MJ s ⁻¹
Baseline SMR plant ref. 49	870	20.1	97.04	81.69
Novel ISEAM process	500	0	108.6	105.9

cooling and a knock-out drum (KOD). A hydrogen purity of 99.99% is maintained throughout regardless of the upstream upset or reaction extent by the use of a Pressure Swing Adsorption (PSA) unit for both the baseline SMR and novel process.

$$\text{Methane conversion}(\%) = \frac{(n_{\text{CH}_4,\text{IN}} - n_{\text{CH}_4,\text{OUT}})}{n_{\text{CH}_4,\text{IN}}} \times 100 \quad (6)$$

$$\% \text{Hydrogen yield} = \frac{\text{Actual hydrogen prod.}}{\text{Theoretical hydrogen prod.}} \times 100 \quad (7)$$

$$\text{Process thermal eff.}(\%) = \frac{(\text{Molar flow of H}_2,\text{out} \times \text{HHV}_{\text{H}_2})}{\text{Molar flow of CH}_4,\text{IN} \times \text{HHV}_{\text{CH}_4} + (\text{net heating utility})} \times 100 \quad (8)$$

$$\text{Hydrogen eff.}(\%) = \frac{(\text{molar flow of H}_2,\text{out} \times \text{HHV}_{\text{H}_2})}{(\text{molar flow of CH}_4,\text{IN} \times \text{HHV}_{\text{CH}_4})} \times 100 \quad (9)$$

3.3 ISEAM process reactor performance

The Carbonator-SMR Reactor is modelled as a packed bed adiabatic catalytic reactor and the membrane reactor is modelled as an iso-thermal catalytic plug flow reactor. Both reactors are characterised with one-dimensional heterogeneous equations. The products profile in the Carbonator-SMR reactor during the pre-breakthrough steady state is shown in Fig. 11. Because of the exothermic carbonation reaction, $(\text{CaO} + \text{CO}_2 \leftrightarrow \text{CaCO}_3, \Delta H_{298}^\circ = -178.8 \text{ kJ mol}^{-1})$, the reactor temperature initially increased to a maximum value of 595 °C within the first 1 m of the reactor length, resulting in faster initial kinetics. The reaction rates become slow afterward as the endothermic reforming dominates. The hydrogen molar fraction rises quickly initially from 19.4% (dry basis) close to the reactor inlet and continues to increase slowly along the reactor length, leaving the Carbonator-SMR reactor at 87% (dry basis). The methane molar fraction drops very quickly initially from 69.5% (dry basis) close to the reactor inlet and continues to decrease slowly along the reactor length, leaving the reactor at 12.7% (dry basis). Fig. 12 shows the Membrane Reactor performance; the reforming and water gas shift reactions are at the slowest rate between the membrane reactor length of 0–8 meters because of the high molar flowrate of hydrogen within these reaction zones (22–87%). The permeation of hydrogen is

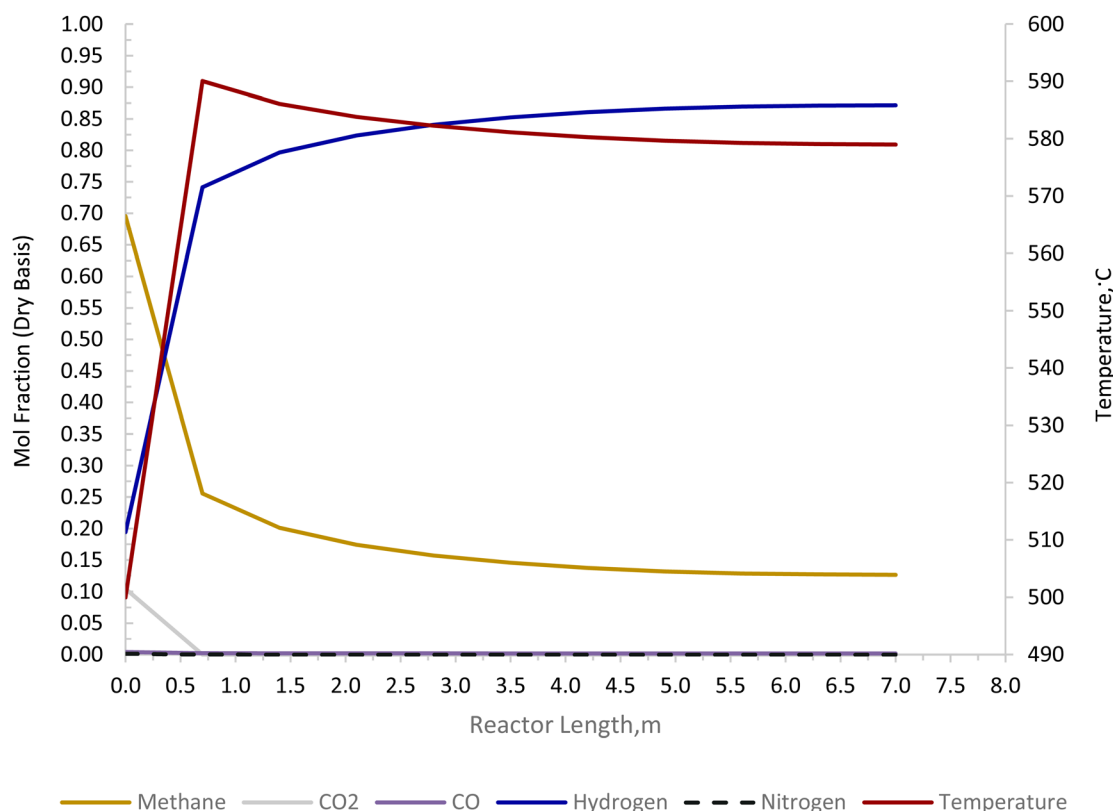


Fig. 11 Carbonator-SMR reactor product profile for the novel ISEAM process (pre-breakthrough steady state).



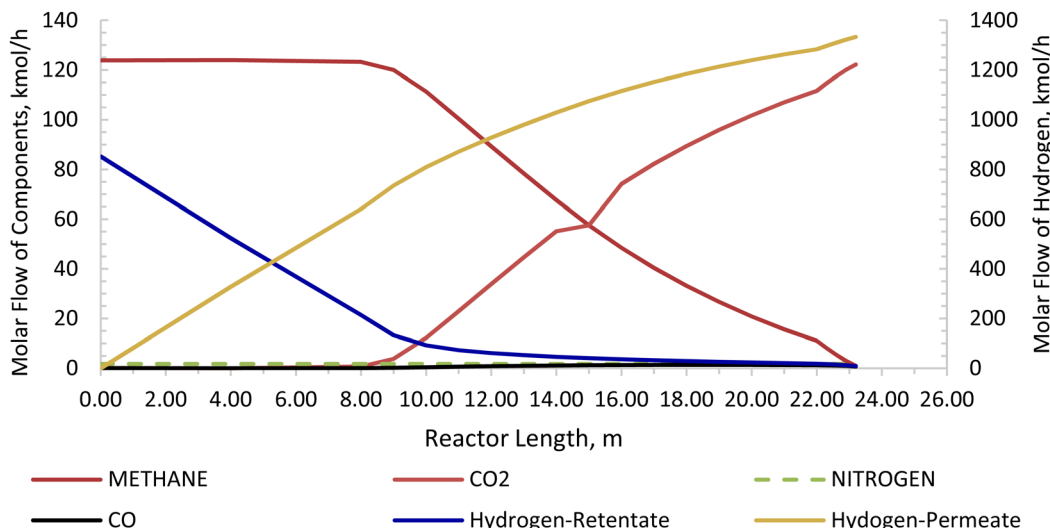


Fig. 12 Membrane reactor product profile for the novel ISEAM process (molar flow of CO_2 , CO , N_2 , H_2 -retentate, and H_2 -permeate).

however at the highest at these locations because of a higher retentate hydrogen partial pressure. As more hydrogen is removed *in situ* from the reaction zones along the membrane reactor, this leads to more methane conversion (and a reduction of methane molar flowrate) and more hydrogen production as per Le-Chatelier's principle. Hydrogen permeation is identified as being as critical as hydrogen selectivity in the design of the Membrane Reactor as more than an average of 54.5% of produced hydrogen was required to be removed from the reactor reaction zones to break the thermochemical reversible equilibrium constraints and shift the reforming reactions to the forward direction.

3.4 Parametric investigations of the novel ISEAM process

Thermodynamic investigations were conducted to determine the optimal operating conditions for the novel process as shown in Fig. 13. A molar ratio of steam to methane is used to control carbon deposition and to ensure forward reactions of steam methane reforming. A steam to methane ratio between 1 and 5 was investigated and Fig. 13(a) shows that an S/C ratio of 3.75–5 gives maximum methane conversion and hydrogen yield. Purge gas is used in the Membrane Reactor to enhance the driving force and to reduce the hydrogen partial pressure in the permeate side, consequently increasing hydrogen permeation as shown in Fig. 13(b) and (c) where the methane conversion and hydrogen yield increases with increasing purge gas (sweep gas). The overall steam methane reforming reaction ($\text{CH}_4 + 2\text{H}_2\text{O} \leftrightarrow \text{CO}_2 + 4\text{H}_2$, $\Delta H_{298}^\circ = 165 \text{ kJ mol}^{-1}$) is endothermic and is favored by high temperature, as shown in Fig. 13(d). For a Carbonator-SMR Reactor and Membrane SMR Reactor temperature of 550°C , the process achieves the maximum methane conversion and hydrogen yield of 99.9% Table 6.

3.5 Economic analysis

The results of the economic model for the baseline SMR process and novel process are summarised in Tables 7, 8 and

Fig. 14–16. The ISEAM process produced 2985 kg h^{-1} of product hydrogen which is approximately 27.2% higher when compared with the baseline SMR process, because of the higher hydrogen production efficiency using the same feed flowrate. As shown in Table 7 and Fig. 14, the total capital cost (TCR) for the baseline SMR is 32.4% higher than the ISEAM process, the first year operating cost (OPEX) for baseline SMR is approx 27% higher than that for the ISEAM process. The levelised cost of hydrogen (LCOH) was calculated as \$2.6 per kg-H_2 for the baseline SMR plant compared with \$1.6 per kg-H_2 for the novel ISEAM process (37.5% cost reduction in favor of the novel process). The novel ISEAM process was further optimised by operating the process units under optimal conditions as per Fig. 13(a) to (d) (S/C = 3.75, reactor temp. = 550°C , purge gas/C1 ratio = 2), which increased the hydrogen production efficiency further (99.9%) giving a calculated LCOH of \$1.3 per kg-H_2 , (50% cost reduction). Fig. 16 shows that the total cost requirement (TCR), commonly referred to as the capitalised cost (CAPEX), which constitutes between 44% and 46% of the LCOH for the three processes, followed by the variable operating and maintenance cost (VOM) which is between 30% and –32% of the LCOH. Sensitivity analysis on the effect of different fuel costs on the LCOH was carried out as shown in Fig. 15 and this effect is minimal. The cost of CO_2 Removal (CCR) was calculated as \$180 per tonne CO_2 for the baseline SMR process compared with \$76.5 per tonne CO_2 for the baseline SMR plant operating with similar feed flowrates (which is a 57.5% cost reduction) in favor of the novel process. The optimized novel process has a Cost of CO_2 Removal (CCR) of \$33.2 per tonne CO_2 (81.6% cost reduction compared to the baseline).

4. Conclusions

A novel blue hydrogen process, the Industrial Sorption Enhanced Auto-Thermal Membrane (ISEAM) process has been developed, modelled, and validated using experimental data from the literature and with thermodynamic equilibrium data.



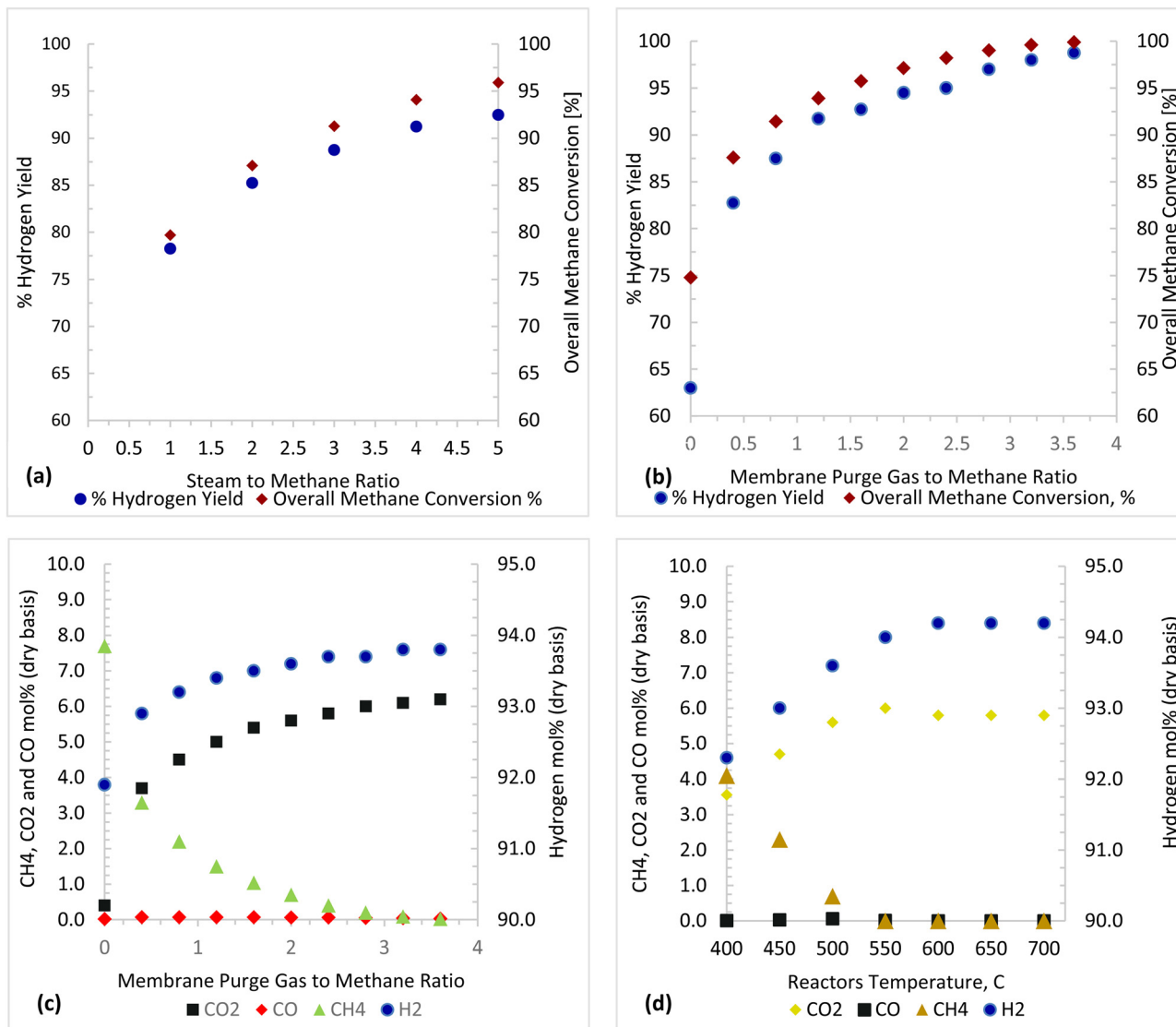


Fig. 13 Parametric investigation results (a) purge gas/CH₄ = 2, reactor temperature: 500 °C (b) and (c) S/C = 3.75, reactor temperature: 500 °C, and (d) purge gas/CH₄ = 2, S/C = 3.75.

Table 7 Main results for economic analysis

Equipment	Baseline SMR	Novel ISEAM process	Optimised novel process
Total bare equipment cost (BEC), \$1000	86 494	63 024	79 568
Total EPCC, \$1000	6918.5	5042	6365
Total process contingency, \$1000	14 930	13 934.9	17 433.8
Total project contingency, \$1000	10 684	8100	10 337
Balance of plant (BOP), \$1000	14 577	10 779	14 486
Total plant cost (TPC), \$1000	133 603.5	100 880	128 190
Owner's cost, \$1000	26 988	20 378	25 894
Total overnight cost (TOC), \$1000	160 591.5	121 258	154 084
Total capital requirement (TCR) or (CAPEX)	183 074	138 234	175 656
First year OPEX	26 766	21 082	26 443
Total hydrogen produced, kg h ⁻¹	2346	2985	4402
First year LCOH	\$2.6 per kg-H ₂	\$1.6 per kg-H ₂	\$1.3 per kg-H ₂
Cost of CO ₂ avoided (CCA) \$ per tonneCO ₂ (zero tax credit)	\$188 per tonneCO ₂	\$65.2 per tonneCO ₂	\$28.2 per tonneCO ₂
Cost of CO ₂ Removal (CCR) \$ per tonneCO ₂ (zero tax credit)	\$180 per tonneCO ₂	\$76.5 per tonneCO ₂	\$33.2 per tonneCO ₂

Table 8 Bare equipment cost results for economic analysis

Equipment	Baseline SMR	Novel ISEAM process	Optimised novel process
Equipment	BEC, \$1000	BEC, \$1000	BEC, \$1000
Feed to plant	315 kmol h ⁻¹ natural gas	315 kmol h ⁻¹ natural gas	500 kmol h ⁻¹ natural gas
	909 kmol h ⁻¹ steam	909 kmol h ⁻¹ steam	1875 kmol h ⁻¹ steam
Hydrogen produced	23.4 MMSCFD (2346 kg h ⁻¹)	29.7 MMSCFD (2985 kg h ⁻¹)	43.8 MMSCFD (4,402 kg h ⁻¹)
SMR Reactor	17 589	—	—
Sulfur polisher (ZnO)	73.895	85.26	107.6
Primary air compressor	2018	1699	2145
Water gas shift reactor	3928	—	—
MDEA CO ₂ unit (95% CO ₂ removal)	25 541	—	—
MEA CO ₂ unit (90% CO ₂) removal	24 228	—	—
PSA unit	11 570	13 349	16 853
Boiler package	1546	1861	2349.5
Carbonator reactor	—	40 773	51475.8
Membrane reactor	—	5257	6637

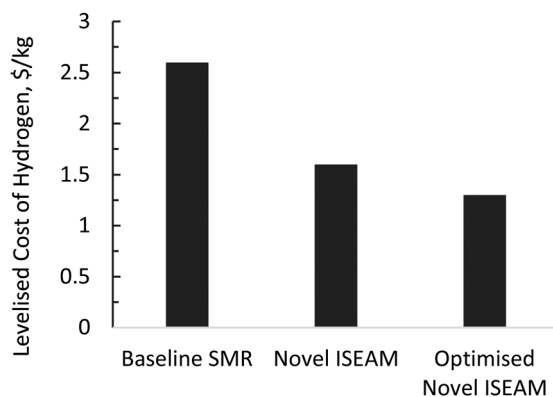


Fig. 14 Levelised cost of hydrogen comparison between baseline SMR process and novel ISEAM process at different fuel cost.

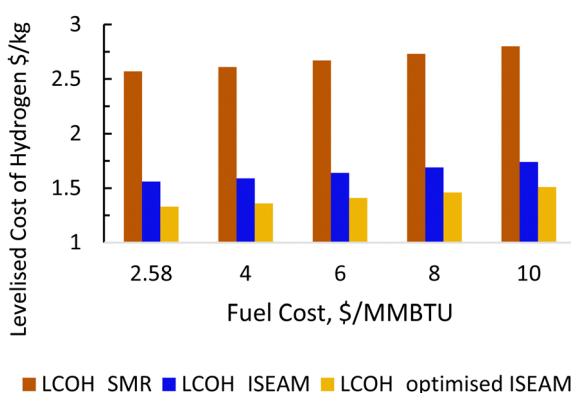


Fig. 15 Economic evaluation comparison at different fuel cost.

The novel process was benchmarked against a baseline Industrial Steam Methane Reforming plant using the same feed compositions and flowrate. The novel process shows improvements in methane conversion, and a hydrogen yield of over 32%. The novel process thermal efficiency of 97.5% (41% higher than the baseline process) is among the highest

compared with alternative processes. Pinch analysis on the novel process shows that the process is auto thermal (no external heating required). Economic evaluation shows a levelized cost of hydrogen (LCOH) of \$1.6 per kg-H₂ and the cost of CO₂ removal (CCR) of \$76.5 per tonneCO₂ for the novel ISEAM process compared to a LCOH of \$2.6 per kg-H₂ and the cost of CO₂ removal (CCR) of \$180 per tonneCO₂ for the baseline SMR plant operating with similar feed flowrates. The optimized novel ISEAM process (at a higher feed rate and higher hydrogen production) gives a LCOH of \$1.3 per kg-H₂ and a cost of CO₂ removal (CCR) of \$33.2 per tonneCO₂. Hydrogen permeation was identified as being as important as hydrogen selectivity in the design of the Membrane Reactor as more than an average of 54.5% of produced hydrogen was required to be removed from the reactor reaction zones to break the thermochemical reversible equilibrium constraints and shift the reforming reactions to the forward direction. The amounts of CO leaving the reactors were reduced to ppm levels. Parametric equilibrium investigations on the novel ISEAM process show that a higher steam-to-methane molar ratio, higher membrane reactor purge gas-to-methane molar ratio, and a higher reactor temperature all increase the methane conversion and hydrogen yield. Optimal operating parameters for the novel ISEAM process were identified. The novel ISEAM process eliminates some of the issues associated with the baseline SMR process such as thermodynamic equilibrium constraints, large amount of utility fuel usage and the use of additional unit operations such as an amine unit to capture the produced CO₂. In addition, steady state operation can be maintained by automating the switching between the sorption/calciation process of the Carbonator-SMR Reactor. The novel ISEAM base process uses a palladium-alloy catalytic membrane reactor and CaO-based Carbonator-SMR Reactor. The capture capacities of CaO-based sorbents are known to decrease rapidly during cycles of carbonation/calciation due to sintering and attrition. Various research works ranging from doping, thermal-pre-treatment, incorporation of inert supports and chemical treatment are being carried out to address the issue of reduced sorption capacity including the use of alternative sorbents such as hydrotalcite. The use of other high temperature membranes such as silica, zeolite, etc,



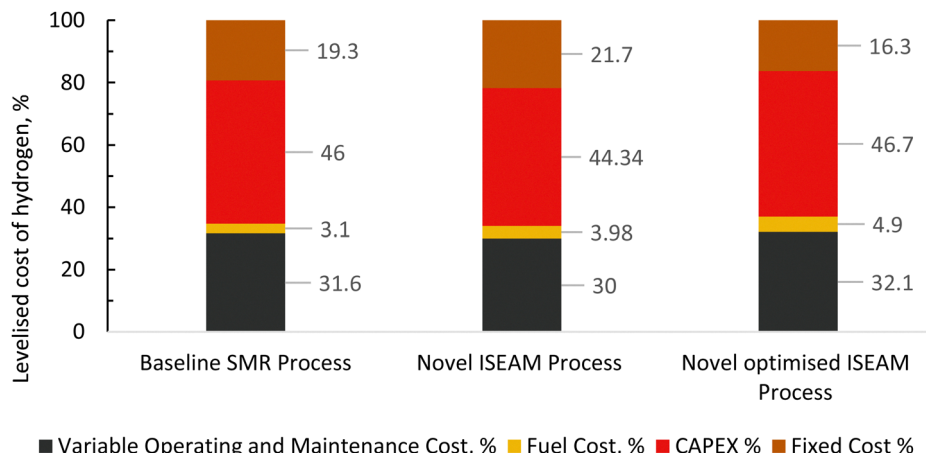


Fig. 16 Distribution of different components of levelized cost (%) for the baseline SMR Process, novel ISEAM and optimised ISEAM process at a fuel cost of \$2.58/MMBTU.

are also being researched. The novel ISEAM process is adaptable and will be able to use such new materials in the future.

Appendix A

Table 9 Experimental data used for carbonator-SMR reactor model validation⁶²

Description	Value
Methane flowrate, NL/h	11.2
Steam: methane ratio	3
Reactor temperature, °C	700
Reactor pressure, kPa	300
GHSV, h ⁻¹	341.5
Bed void fraction	0.5
CaO density, kg m ⁻³	1257
Reactor diameter, m	0.024
Reactor length, m	0.29

Conflicts of interest

There are no conflicts to declare.

Table 10 Experimental data used for membrane reactor model validation⁴⁴

Parameters	Values	Units
Membrane diameter	0.0127	m
Length of membrane tube	0.25	m
Membrane thickness	6	µm
Methane feed flow	0.78	mol h ⁻¹
Membrane sweep gas flowrate	0.5	mol h ⁻¹
Reactor pressure	350	kPa
Reactor temperature	500	°C
Retentate pressure	100	kPa
Membrane permeance	0.00346	mol m ⁻² s ⁻¹ Pa ^{0.5}





Appendix B

Table 11 Rate and kinetic equations for the main reactions

Reactions	Rate equations	Kinetic equation and constants	Ref.
$\text{CH}_4 + \text{H}_2\text{O} \rightarrow \text{CO} + 3\text{H}_2, \quad \Delta H_{298}^\circ = 206 \text{ kJ mol}^{-1}$	$R_1 = \frac{k_1 \left(\frac{P_{\text{CH}_4} P_{\text{H}_2\text{O}}}{P_{\text{H}_2}^{2.5}} - P_{\text{H}_2} 0.5 P_{\text{CO}} / K_1 \right)}{\text{DEN}^2}$	(10) $k_1 = 1.17 \times 10^{12} \exp(-240 \text{ 100/RT})$	(13) 4,40,51,52
$\text{CO} + \text{H}_2\text{O} \leftrightarrow \text{CO}_2 + \text{H}_2, \quad \Delta H_{298}^\circ = -41 \text{ kJ mol}^{-1}$	$R_2 = \frac{k_2 \left(\frac{P_{\text{CO}} P_{\text{H}_2\text{O}}}{P_{\text{H}_2}} - P_{\text{CO}_2} / K_2 \right)}{\text{DEN}^2}$	(11) $k_2 = 5.42 \times 10^{12} \exp(67130/RT)$	(14) 4,40,51,52
$\text{CH}_4 + 2\text{H}_2\text{O} \leftrightarrow \text{CO}_2 + 4\text{H}_2, \quad \Delta H_{298}^\circ = 165 \text{ kJ mol}^{-1}$	$R_3 = \frac{k_3 \left(\frac{P_{\text{CH}_4} P_{\text{H}_2\text{O}}^2}{P_{\text{H}_2}^{3.5}} - P_{\text{H}_2} 0.5 P_{\text{CO}_2} / K_1 K_2 \right)}{\text{DEN}^2}$	(12) $k_3 = 2.83 \times 10^{11} \exp(-243900/RT)$	(15) 4,40,51,52
$\text{CaO} + \text{CO}_2 \leftrightarrow \text{CaCO}_3, \quad \Delta H_{298}^\circ = -178.8 \text{ kJ mol}^{-1}$		$\text{DEN} = 1 + K_{\text{CO}} P_{\text{CO}} + K_{\text{H}_2} P_{\text{H}_2} + K_{\text{CH}_4} P_{\text{CH}_4} + K_{\text{H}_2\text{O}} P_{\text{H}_2\text{O}}$	(16) 4,40,51,52
		$k_{\text{CH}_4} = 6.65 \times 10^{-4} \exp(38280/RT)$ $k_{\text{H}_2\text{O}} = 1.77 \times 10^5 \exp(-88680/RT)$ $k_{\text{CO}} = 8.25 \times 10^{-5} \exp(70650/RT)$ $k_{\text{H}_2} = 6.15 \times 10^{-9} \exp(82900/RT)$ $k_1 = 4.71 \times 10^{12} \exp(-324000/RT)$ $k_2 = 1.14 \times 10^{-12} \exp(37300/RT)$ $K_3 = K_1 K_2$	(17) 4,40,51,52 (18) 4,40,51,52 (19) 4,40,51,52 (20) 4,40,51,52 (21) 4,40,51,52 (22) 4,40,51,52 (23) 4,40,51,52
		$\frac{dX}{dt} = \phi_{\text{f}} k_{\text{reac-max}} X_{\text{active}} (V_{\text{CO}_2} - V_{\text{CO}_2,\text{eq}})$ $k_{\text{reac-max}} = 0.36 \text{ s}^{-1}$ $V_{\text{CO}_2,\text{eq}} = 4.137 \times 10^7 \exp\left(\frac{-20474}{T}\right)$	(24) 40,53 (25) 40,53 (26) 40,53
		The hydrogen flow from the retentate to the permeate side of the membrane reactor is governed by Sievert's law: $F_{\text{H}_2,\text{permeance}} = H_{2\text{permeance}} \times \text{Area} \times \chi \sqrt{P_{\text{H}_2,\text{ret}} - P_{\text{H}_2,\text{eq}}}$	(27) 40,44

Notes and references

1 US Energy Information Administration, dated 13th April, 2021.

2 Hydrogen scaling up. A sustainable pathway for the global energy transition, Hydrogen Council (dated 2017), (www.hydrogencouncil.com/wpcontent/uploads/2017/11/hydrogen-scaling-up-hydrogencouncil.pdf).

3 D. Celik and M. Yildiz, Investigation of hydrogen production methods in accordance with green chemistry principles, *Int. J. Hydrogen Energy*, 2017, **42**(36), 23395–23401.

4 H. Lee, B. Lee, M. Byun and H. Lim, Comparative techno-economic analysis for steam methane reforming in a sorption-enhanced membrane reactor: Simultaneous H₂ production and CO₂ capture, *Chem. Eng. Res. Des.*, 2021, **171**, 383–394.

5 Y. Yan, V. Manovic, E. J. Anthony and P. T. Clough, Techno-economic analysis of low-carbon hydrogen production by sorption enhanced steam methane reforming (SE-SMR) processes, *Energy Convers. Manage.*, 2020, **226**, 113530.

6 P. Nikolaidis and A. Poullikkas, A comparative overview of hydrogen production processes, *Renewable Sustainable Energy Rev.*, 2017, **67**, 597–611.

7 L. Zhu, L. Li and J. Fan, A modified process for overcoming the drawbacks of conventional steam methane reforming for hydrogen production: Thermodynamic investigation, *Chem. Eng. Res. Des.*, 2015, **104**, 792–806.

8 Y. Ma, G. Guan, X. Hao, J. Cao and A. Abudula, Molybdenum carbide as alternative catalyst for hydrogen production – A review, *Renewable Sustainable Energy Rev.*, 2017, **75**, 1101–1129.

9 C.-H. Chen, C.-T. Yu and W.-H. Chen, Improvement of steam methane reforming via in-situ CO₂ sorption over a nickel-calcium composite catalyst, *Int. J. Hydrogen Energy*, 2021, **46**(31), 16655–16666.

10 A. Di Giuliano and K. Gallucci, Sorption enhanced steam methane reforming based on nickel and calcium looping: a review, *Chem. Eng. Process.*, 2018, **130**, 240–252.

11 S. Z. Abbas, V. Dupont and T. Mahmud, Modelling of high purity H₂ production via sorption enhanced chemical looping steam reforming of methane in a packed bed reactor, *Fuel (Guildford)*, 2017, **202**, 271–286.

12 A. Phluanglue, W. Khaodee and S. Assabumrungrat, Simulation of intensified process of sorption enhanced chemical-looping reforming of methane: Comparison with conventional processes, *Comput. Chem. Eng.*, 2017, **105**, 237–245.

13 S. Abuelgasim, W. Wang and A. Abdalazeez, A brief review for chemical looping combustion as a promising CO₂ capture technology: Fundamentals and progress, *Sci. Total Environ.*, 2021, **764**, 142892.

14 S. Z. Abbas, V. Dupont and T. Mahmud, Modelling of H₂ production in a packed bed reactor via sorption enhanced steam methane reforming process, *Int. J. Hydrogen Energy*, 2017, **42**(30), 18910–18921.

15 J. R. Fernandez, J. C. Abanades and R. Murillo, Modeling of sorption enhanced steam methane reforming in an adiabatic fixed bed reactor, *Chem. Eng. Sci.*, 2012, **84**, 1–11.

16 J. R. Fernandez, J. C. Abanades and G. Grasa, Modeling of sorption enhanced steam methane reforming—Part II: Simulation within a novel Ca/Cu chemical loop process for hydrogen production, *Chem. Eng. Sci.*, 2012, **84**, 12–20.

17 G. Ji, J. G. Yao, P. T. Clough, J. C. D. da Costa, E. J. Anthony and P. S. Fennell, *et al.*, Enhanced hydrogen production from thermochemical processes, *Energy Environ. Sci.*, 2018, **11**(1), 2647–2672.

18 M. Broda, V. Manovic, Q. Imtiaz, A. M. Kierzkowska, E. J. Anthony and C. R. Müller, High-Purity Hydrogen via the Sorption-Enhanced Steam Methane Reforming Reaction over a Synthetic CaO-Based Sorbent and a Ni Catalyst, *Environ. Sci. Technol.*, 2013, **47**(11), 6007–6014.

19 J. R. Fernández and J. C. Abanades, Sorption enhanced reforming of methane combined with an iron oxide chemical loop for the production of hydrogen with CO₂ capture: Conceptual design and operation strategy, *Appl. Therm. Eng.*, 2017, **125**, 811–822.

20 M. Erans, V. Manovic and E. J. Anthony, Calcium looping sorbents for CO₂ capture, *Appl. Energy*, 2016, **180**, 722–742.

21 B. Dou, Y. Song, Y. Liu and C. Feng, High temperature CO₂ capture using calcium oxide sorbent in a fixed-bed reactor, *J. Hazard. Mater.*, 2010, **183**(1–3), 759–765.

22 I. Aloisi, A. Di Giuliano, A. Di Carlo, P. U. Foscolo, C. Courson and K. Gallucci, Sorption enhanced catalytic Steam Methane Reforming: Experimental data and simulations describing the behaviour of bi-functional particles, *Chem. Eng. J.*, 2017, **314**, 570–582.

23 I. Aloisi, N. Jand, S. Stendardo and P. U. Foscolo, Hydrogen by sorption enhanced methane reforming: A grain model to study the behavior of bi-functional sorbent-catalyst particles, *Chem. Eng. Sci.*, 2016, **149**, 22–34.

24 J. Adanez, A. Abad, F. Garcia-Labiano, P. Gayan and L. F. de Diego, Progress in Chemical-Looping Combustion and Reforming technologies, *Prog. Energy Combust. Sci.*, 2012, **38**(2), 215–282.

25 J. Boon, V. Spallina, Y. van Delft and M. van Sint Annaland, Comparison of the efficiency of carbon dioxide capture by sorption-enhanced water-gas shift and palladium-based membranes for power and hydrogen production, *Int. J. Greenhouse Gas Control*, 2016, **50**, 121–134.

26 F. H. Alshafei, L. T. Minardi, D. Rosales, G. Chen and A. Simonetti Dante, Improved Sorption-Enhanced Steam Methane Reforming via Calcium Oxide-Based Sorbents with Targeted Morphology, *Energy Technol.*, 2019, **7**(3), 1800807.

27 H. Yu, H. Sun, G. Bao, H. Liu, J. Hu and H. Wang, Computational fluid dynamics study of hydrogen production by sorption enhanced steam ethanol reforming process in fluidized bed, *Fuel (Guildford)*, 2023, **344**, 128043.

28 C. Udemu and C. Font-Palma, Modelling of sorption-enhanced steam reforming (SE-SR) process in fluidised bed reactors for low-carbon hydrogen production: A review, *Fuel (Guildford)*, 2023, **340**, 127588.

29 A. Mostafa, I. Rapone, A. Bosetti, M. C. Romano, A. Beretta and G. Groppi, Modelling of methane sorption enhanced reforming for blue hydrogen production in an adiabatic



fixed bed reactor: unravelling the role of the reactor's thermal behavior, *Int. J. Hydrogen Energy*, 2023, **48**, 26475–26491.

30 B. Lee and H. Lim, Cost-competitive methane steam reforming in a membrane reactor for H₂ production: Technical and economic evaluation with a window of a H₂ selectivity, *Int. J. Energy Res.*, 2019, **43**(4), 1468–1478.

31 A. Iulianelli, S. Liguori, J. Wilcox and A. Basile, Advances on methane steam reforming to produce hydrogen through membrane reactors technology: A review, *Catal. Rev.: Sci. Eng.*, 2016, **58**(1), 1–35.

32 X. Wu, C. Wu and S. Wu, Dual-enhanced steam methane reforming by membrane separation of H₂ and reactive sorption of CO₂, *Chem. Eng. Res. Des.*, 2015, **96**, 150–157.

33 M. A. Murmura, S. Cerbelli and M. C. Annesini, An equilibrium theory for catalytic steam reforming in membrane reactors, *Chem. Eng. Sci.*, 2017, **160**, 291–303.

34 H. Lee, A. Kim, B. Lee and H. Lim, Comparative numerical analysis for an efficient hydrogen production via a steam methane reforming with a packed-bed reactor, a membrane reactor, and a sorption-enhanced membrane reactor, *Energy Convers. Manage.*, 2020, **213**, 112839.

35 K. Ghasemzadeh, R. Zeynali, A. Basile and A. Iulianelli, CFD analysis of a hybrid sorption-enhanced membrane reactor for hydrogen production during WGS reaction, *Int. J. Hydrogen Energy*, 2017, **42**(43), 26914–26923.

36 J. M. Alarcón, J. R. Fernández and J. C. Abanades, Study of a Cu-CuO chemical loop for the calcination of CaCO₃ in a fixed bed reactor, *Chem. Eng. J.*, 2017, **325**, 208–220.

37 J. R. Fernández and J. C. Abanades, Overview of the Ca–Cu looping process for hydrogen production and/or power generation, *Curr. Opin. Chem. Eng.*, 2017, **17**, 1–8.

38 J. C. Abanades, R. Murillo, J. R. Fernandez, G. Grasa and I. Martínez, New CO₂ Capture Process for Hydrogen Production Combining Ca and Cu Chemical Loops, *Environ. Sci. Technol.*, 2010, **44**(17), 6901–6904.

39 L. Díez-Martín, I. Martínez, G. Grasa and R. Murillo, Investigation of the reduction kinetics of high loaded CuO-based materials suitable for the Ca/Cu looping process, *Fuel (Guildford)*, 2018, **230**, 376–389.

40 M. M. Shahid, S. Z. Abbas, F. Maqbool, S. Ramirez-Solis, V. Dupont and T. Mahmud, Modeling of sorption enhanced steam methane reforming in an adiabatic packed bed reactor using various CO₂ sorbents, *J. Environ. Chem. Eng.*, 2021, **9**(5), 105863.

41 G. Diglio, D. P. Hanak, P. Bareschino, F. Pepe, F. Montagnaro and V. Manovic, Modelling of sorption-enhanced steam methane reforming in a fixed bed reactor network integrated with fuel cell, *Appl. Energy*, 2018, **210**, 1–15.

42 S. M. Nazir, J. H. Cloete, S. Cloete and S. Amini, Pathways to low-cost clean hydrogen production with gas switching reforming, *Int. J. Hydrogen Energy*, 2021, **46**(38), 20142–20158.

43 J. D. Silva and C. A. M. de Abreu, Modelling and simulation in conventional fixed-bed and fixed-bed membrane reactors for the steam reforming of methane, *Int. J. Hydrogen Energy*, 2016, **41**(27), 11660–11674.

44 C.-H. Kim, J.-Y. Han, H. Lim, D.-W. Kim and S.-K. Ryu, Methane steam reforming in a membrane reactor using high-permeable and low-selective Pd-Ru membrane, *Korean J. Chem. Eng.*, 2017, **34**(4), 1260–1265.

45 G. Ye, D. Xie, W. Qiao, J. R. Grace and C. J. Lim, Modeling of fluidized bed membrane reactors for hydrogen production from steam methane reforming with Aspen Plus, *Int. J. Hydrogen Energy*, 2009, **34**(11), 4755–4762.

46 A. Shafiee, M. Arab, Z. Lai, Z. Liu and A. Abbas, Modelling and sequential simulation of multi-tubular metallic membrane and techno-economics of a hydrogen production process employing thin-layer membrane reactor, *Int. J. Hydrogen Energy*, 2016, **41**(42), 19081–19097.

47 G. Ji, M. Zhao and G. Wang, Computational fluid dynamic simulation of a sorption-enhanced palladium membrane reactor for enhancing hydrogen production from methane steam reforming, *Energy (Oxford)*, 2018, **147**, 884–895.

48 B. Dou, K. Wu, H. Zhang, B. Chen, H. Chen and Y. Xu, Sorption-enhanced chemical looping steam reforming of glycerol with CO₂ in-situ capture and utilization, *Chem. Eng. J.*, 2023, **452**, 139703.

49 M. Salem, A. M. Shoaib and A. F. M. Ibrahim, Simulation of a Natural Gas Steam Reforming Plant for Hydrogen Production Optimization, *Chem. Eng. Technol.*, 2021, **44**(9), 1651–1659.

50 M. Salem, A. M. Shoaib and A. F. M. Ibrahim, Simulation of a Natural Gas Steam Reforming Plant for Hydrogen Production Optimization, *Chem. Eng. Technol.*, 2021, **44**(9), 1651–1659.

51 W.-J. Sheu, C.-Y. Chang and Y.-C. Chen, Transient reaction phenomena of sorption-enhanced steam methane reforming in a fixed-bed reactor, *Int. J. Hydrogen Energy*, 2022, **47**(7), 4357–4374.

52 J. Xu and G. F. Froment, Methane steam reforming, methanation and water-gas shift: I. Intrinsic kinetics, *AIChE J.*, 1989, **35**(1), 88–96.

53 N. Rodriguez, M. Alonso and J. C. Abanades, Experimental investigation of a circulating fluidized-bed reactor to capture CO₂ with CaO, *AIChE J.*, 2011, **57**(5), 1356–1366.

54 GCCSI. Global CCS Institute-toward a common method of cost estimation for CO₂ capture and storage at fossil fuel power plants. 2013.

55 Oxera. Discount rates for low-carbon and renewable generation technologies Prepared for the Committee on Climate Change. 2011.

56 M. N. Khan and T. Shamim, Techno-economic assessment of a plant based on a three reactor chemical looping reforming system, *Int. J. Hydrogen Energy*, 2016, **41**(48), 22677–22688.

57 G. Collodi, G. Azzaro, N. Ferrari and S. Santos, Techno-economic Evaluation of Deploying CCS in SMR Based Merchant H₂ Production with NG as Feedstock and Fuel, *Energy Procedia*, 2017, **114**, 2690–2712.

58 E. Callum, *et al.*, Novel Steam Methane/Gas Heated Reformer Phase 1 Final Study Report, Wood Group UK Limited: UK, 2020, p. 22.



59 M. Haaf, R. Anantharaman, S. Roussanaly, J. Ströhle and B. Epple CO₂ capture from waste-to-energy plants: Techno-economic assessment of novel integration concepts of calcium looping technology. 2020.

60 National Energy Technology Laboratory, Assessment of hydrogen Production with CO₂ Capture Volume 1: Baseline-State-Of the-Arts Plant, Revision 1, Nov 14th, 2011, DOE/NETL-2011/1434.

61 Department for Business Energy & Industrial Strategy, Hydrogen Production Costs, 2021.

62 D. K. Lee, I. H. Baek and W. L. Yoon, Modeling and simulation for the methane steam reforming enhanced by in situ CO₂ removal utilizing the CaO carbonation for H₂ production, *Chem. Eng. Sci.*, 2004, **59**(4), 931–942.

63 H. Bahzad, N. Shah, N. M. Dowell, M. Boot-Handford, S. M. Soltani and M. Ho, *et al.*, Development and techno-economic analyses of a novel hydrogen production process via chemical looping, *Int. J. Hydrogen Energy*, 2019, **44**(39), 21251–21263.

64 S. Z. Abbas, V. Dupont and T. Mahmud, Kinetics study and modelling of steam methane reforming process over a NiO/Al₂O₃ catalyst in an adiabatic packed bed reactor, *Int. J. Hydrogen Energy*, 2017, **42**(5), 2889–2903.

