Experimental Characterization and Energy Performance Assessment of a Sorption-Enhanced Steam–Methane Reforming System

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Abstract: The production of blue hydrogen through sorption-enhanced processes has emerged as a suitable option to reduce greenhouse gas emissions. Sorption-enhanced steam–methane reforming (SESMR) is a process intensification of highly endothermic steam–methane reforming (SMR), ensured by in situ carbon capture through a solid sorbent, making hydrogen production efficient and more environmentally sustainable. In this study, a comprehensive energy model of SESMR was developed to carry out a detailed energy characterization of the process, with the aim of filling a current knowledge gap in the literature. The model was applied to a bench-scale multicycle SESMR/sorbent regeneration test to provide an energy insight into the process. Besides the experimental advantages of higher hydrogen concentration (90 mol% dry basis, 70 mol% wet basis) and performance of CO₂ capture, the developed energy model demonstrated that SESMR allows for substantially complete energy self-sufficiency through the process. In comparison to SMR with the same process conditions (650 °C, 1 atm) performed in the same experimental rig, SESMR improved the energy efficiency by about 10%, further reducing energy needs.

Keywords: hydrogen production; sorption-enhanced steam–methane reforming (SESMR); SESMR energy theoretical model; CO₂ capture; CO₂ emissions reduction

1. Introduction

The fight against climate change has returned to the center of the global debate, after wide media coverage of the promulgation of the Paris agreement in December 2015. This agreement was ratified in November 2016 and legally bound its Parties to limit global warming to well below 2 °C, preferably to 1.5 °C, compared to pre-industrial levels. Countries have planned to reach a global peak of greenhouse gas emissions (GHGE) as soon as possible, in order to achieve climate neutrality by mid-century [1].

China, accounting for 28% of global emissions, recently set the target of carbon neutrality achievement for before 2060 [2]. After its establishment in December 2019, the European Commission launched the European Green Deal, a roadmap of legislative and economic measures that pursue EU climate neutrality by 2050 [3,4]; this commitment was confirmed even during the COVID-19 pandemic [5]. In February 2021, the United States officially rejoined the Paris agreement [6]. The decoupling of economic growth and GHGE was demonstrated not only as possible but also as suitable in order to boost the new so-called green economies [7].

In this framework, an effective transition is occurring from the current exploitation of fossil fuels to renewable energy sources or low-carbon technologies. Carbon capture and storage (CCS) coupled with fossil fuel exploitation may play a relevant role in making this transition affordable, as also assessed by the United Nations Intergovernmental Panel on Climate Change (UNIPCC) [8–10].
Inside this framework, in recent years, academic and technological interest has grown regarding sorption enhanced steam methane reforming (SESMR) [11]. This process produces “blue hydrogen”, i.e., H\textsubscript{2} obtained from fossil fuel capturing the related CO\textsubscript{2} emissions [12]. H\textsubscript{2} has gained appeal as an alternative to fossils fuels for several decades [13,14] (the locution “hydrogen economy” was first introduced by General Motors Co. in 1970 [15]), and nowadays, it is considered an interesting energy vector for renewable energy storage as well (the so-called “green hydrogen”) [12]. As a matter of fact, H\textsubscript{2}-based technologies have already begun to spread, with H\textsubscript{2} fueling stations [16] for direct use in internal combustion engines (ICEs)—either pure or blended with fossil fuels—and for feeding fuel cell electric vehicles [17], which appear to be the true new paradigm for sustainable mobility. Stationary energy stations for electricity production based on fuel cell technology are envisaged as well.

Currently, steam methane reforming (SMR) of natural gas by Ni-based catalysts is the main commercial technology to produce H\textsubscript{2}, performed in vertical tubular packed-bed reactors (800–1000 °C, 1.5–3.0 MPa) and heated inside flame furnaces [18–20]. This productive pathway covers nearly half of the world’s H\textsubscript{2} demand [15] and it is technologically mature [21,22]. Nonetheless, the entire SMR industrial process involves some critical issues as far as the current climate situation is concerned: (i) the SMR process is highly endothermic—the reactors intensively require heat, usually provided by burning part of the natural gas feedstock and causing relevant net CO\textsubscript{2} emissions; (ii) the temperature decrease associated with downstream operations of CO\textsubscript{2} removal (e.g., shift steps or amine scrubbing) determines poor energy integration efficiency [23].

SESMR, which has not yet been developed at the industrial scale [11], may represent an affordable and more sustainable route to help the transition towards a hydrogen economy in agreement with the UNIPCC suggestions, as it couples the SMR (Equation (1)) process with the in situ carbon capture operated by a solid CO\textsubscript{2} sorbent, therefore intensifying the process of traditional SMR [23–25]. Besides CO\textsubscript{2} capture, SESMR brings an additional advantage—in situ CO\textsubscript{2} removal from the gaseous phase makes water-gas shift (WGS, Equation (2)) and, therefore, SMR equilibria move towards products, that is, to further H\textsubscript{2} production. As a result, high-purity H\textsubscript{2} can be obtained by an SESMR process [26,27]. Among high-temperature solid sorbents for SESMR, CaO is a very typical choice, due to its wide availability in low-cost natural mineralized forms (e.g., limestone or dolomite), its high stoichiometric sorption capacity (78.6 g CO\textsubscript{2} per 100 g CaO), and its fast carbonation (CBN, Equation (3)) kinetics in the temperature window compatible with SMR [28].

\[
\begin{align*}
\text{CH}_4(\text{g}) + H_2O(\text{v}) &\leftrightarrow CO(\text{g}) + 3H_2(\text{g}) \quad \Delta H_{R298}^0 = 206.2 \text{ kJ mol}^{-1} \quad (1) \\
CO(\text{g}) + H_2O(\text{v}) &\leftrightarrow CO_2(\text{g}) + H_2(\text{g}) \quad \Delta H_{R298}^0 = -41.2 \text{ kJ mol}^{-1} \quad (2) \\
CO_2(\text{g}) + CaO(\text{s}) &\leftrightarrow CaCO_3(\text{s}) \quad \Delta H_{R298}^0 = -178.2 \text{ kJ mol}^{-1} \quad (3)
\end{align*}
\]

With CaO as the sorbent and Ni as the catalyst, 1 atm and 650 °C are good process conditions for SESMR, milder than those of traditional SMR [29,30]. The solid CO\textsubscript{2} sorbent eventually gets saturated, and then, continuous H\textsubscript{2} production requires a cyclical alternation between suitable process conditions for SESMR and those for sorbent regeneration, the latter depending on the CBN equilibrium [31]. With regard to this alternation of conditions from the solids’ point of view, the scientific literature mainly proposes two reactor configurations: (i) packed bed reactor couples [32,33], or (ii) dual fluidized bed reactor systems with internal recirculation of bed particles [34]. With both, one reactor works as a reformer and the other as a calciner for sorbent regeneration.

SESMR can be performed by raw mixing of the solo-catalyst and the solo-sorbent particles (two-particle system) [35–37], or by combined sorbent–catalyst materials (CSCMs), that is, solids containing both CO\textsubscript{2} sorbent and SMR catalyst phases [38–43]. Compared to two-particle systems, CSCMs allow for a lower solid hold-up, ensuring a more intimate
compensation between endothermic and exothermic reactions, and decreasing inter-particle and intra-particle resistances to mass transfer [44–49].

From an energy point of view, the heat required during the endothermic reforming processes, usually provided by an external furnace, is certainly one of the main drawbacks of this technology [32]. When SESMR as a continuous operation is considered, the need to cyclically regenerate the sorbent is presented as an additional significant issue by a temperature swing, which triggers endothermic calcination (reverse of CBN, (3)) [50]. In the literature, it has been proposed to provide the energy required for this step by direct combustion of the fuel in an oxy-fuel reactor [51,52], thus increasing the environmental impact of the produced hydrogen.

To fully cover the energy demands of the whole SESMR process, ensuring complete energy self-sufficiency, a thermodynamic characterization is needed, as done in [53] for steam, dry, and autothermal methane reforming.

As a matter of fact, the combination in one reactor vessel of Reactions (1)–(3) due to SESMR (i.e., integrating H₂-producing reactions and CO₂ separation) achieves higher hydrogen yield [54–56]. In addition, it potentially enables the compensation between the energy required by SMR (1) and heat released by WGS (2) and CO₂ capture by CaO (3) [57].

To define a complete thermal integration, comprehensive modeling of an SESMR plant was performed in [50]. A complete thermal integration was achieved between the available hot stream in the plant, leading to the production of power through high-pressure steam. In [32], the feasibility of the SESMR process through a network of packed bed reactors was assessed, and the design and operating parameters under energy self-sufficiency conditions were defined. In [33], a thermo-economic analysis of the proposed solution was performed, and it was found that both SESMR and SESMR integrated with a solid oxide fuel cell achieve higher efficiency than that from the reference SMR case.

In the bench-scale packed bed reactor developed in [58], the feasibility of an innovative SESMR process was experimentally demonstrated, where the heat required by the endothermic sorbent regeneration stage was given by the second loop of NiO-based oxygen carrier.

Regarding the energy aspects related to the process configuration, the adoption of fluidized bed reactors ensures SESMR performs continuously, achieving great mass- and heat-transfer characteristics, but usually at low pressure [23,59–61]. To easily operate at high pressure and avoid gas/solid separation from a heterogeneous stream, packed-bed architecture was recently proposed as a reliable alternative solution [61,62]. Nevertheless, the packed-bed configuration involves restrictions in terms of sorbent regeneration and limited heat-transfer, and, thus, sorbent–catalyst separation is generally proposed for small-scale production [63].

In the framework of the European research project ASCENT (European Union’s Seventh Framework Program grant agreement No. 608512), several academic and industrial partners investigated SESMR feasibility until the level of a robust proof of concept at the pilot-scale, with a system based on fluidized beds. The University of L’Aquila, in cooperation with the University of Strasbourg, developed several CSCMs based on Ni-CaO-Ca₁₂Al₁₄O₃₃ for SESMR, from both experimental and modeling points of view [48,49,64–71].

In this work, the case of study of SESMR performed by the CSCM with 10 wt% of Ni was investigated, obtained by wet impregnation with Ni-nitrate of a CaO-based sorbent with 15 wt% of free-CaO on mayenite (Ca₁₂Al₁₄O₃₃). This CSCM, hereafter referred to as CaO15Ni10, was developed for multicycle SESMR/sorbent regeneration and tested in a packed bed reactor rig at a bench-scale [63,67].

The focus was on the energy retrofit of this SESMR application. A comprehensive thermodynamic model of the system was developed in this study based on an experimental bench-scale prototype. Due to the large set of experimental data, the model provided reliable and robust results in terms of the energy characterization of the endothermic and exothermic reactions involved in the different stages of the multicycle SESMR/sorbent-
regeneration process. In this way, the energy performance of SESMR was quantified, opening the way for optimization of the SESMR process in future studies.

2. Materials and Methods

2.1. SESMR Experiments

CaO15Ni10 was tested in a bench-scale packed-bed reactor facility, performing multicycle SESMR/sorbent regeneration tests. The experimental apparatus is fully detailed elsewhere [51,53], but for the sake of clarity it is also described here. A stainless steel vertical tubular reactor hosted the CSCM packed bed (nominal outside diameter = \( \frac{3}{4} \)”, 6.8 g of CSCM), heated by an electric furnace. SESMR was performed at 650 °C and 1 atm. The flow rates of inlet gases (H\(_2\), CH\(_4\), CO\(_2\), N\(_2\)) and steam were properly controlled, for example, the weight hourly space velocity—in reference to CH\(_4\) and Ni—equaled 2.40 Nl\(_{\text{CH}_4}\) h\(^{-1}\) g\(_{\text{Ni}}\)\(^{-1}\) during SESMR (the value previously selected for CaO15Ni10 in [65]). Downstream, unreacted water was separated by condensation and the overall molar flow rates of dried products were measured, as well as their compositions (vol% of H\(_2\), CH\(_4\), CO, CO\(_2\)).

Di Giuliano et al. [63] tested CaO15Ni10 throughout 205 consecutive reforming cycles alternating with sorbent–regenerations. Every four or five cycles, a long duration (1 h) was set so that the CSCM sorption capacity was completely saturated. In these cases, the first part of the cycle corresponded to SESMR (1–3), while in its final part—that is, after saturation—only SMR (1) and WGS (2) could occur. For the other cycles, a shorter duration was chosen (10 min), interrupting the process before the occurrence of sorbent saturation, and thus, recording only an SESMR phase.

In this work, the first 80 reforming cycles out of the 205 were used as a reference for the energy study, involving both long and short reforming cycles. For all 80 cycles, the SESMR energy features were evaluated. In addition, for the long cycles, a comparison was performed between the features of reforming in the presence or absence of sorption-enhancing effects.

2.2. Energetic Model of SESMR

A mathematical model was developed to perform an energy retrofit of the SESMR process. Starting from the experimental configuration, the first step was the schematization of the process from an energy point of view. In the model scheme shown in Figure 1, the power contributions of the main stages of the processes can be recognized:

(a) Subcooled water enters the evaporator, where it receives a thermal power \( P_{\text{th1}} \). Due to this thermal exchange, the water leaves the evaporator as superheated vapor (steam) at 300 °C.

(b) The superheated steam is mixed with methane at ambient temperature.

(c) The mixture of methane and superheated steam enters a heat exchanger, receiving the thermal power \( P_{\text{th2}} \) and reaching the state required to be fed into the SESMR reactor.

(d) The feed (methane and steam) enters the reactor at the required condition \( T_{(d)} = 650 \) °C). From the energy point of view, two situations occur:

(1) The SMR endothermic Reaction (1) requires a thermal power \( P_{\text{th3}} \) to be provided to the system. This power also takes into account the slightly exothermic contribution of WGS (2);

(2) The CBN exothermic Reaction (3) makes the thermal power \( P_{\text{th4}} \) available.

(e) The sorbent regeneration at 950 °C requires that a significant amount of thermal power \( P_{\text{th5}} \) is provided to heat up the CSCM at the temperature in correspondence to that with which the calcination takes place.

(f) \( P_{\text{th6}} \) is the power required by the calcination (reverse of Reaction (3)), which is a highly endothermic reaction.

(g) Subsequently, the regenerated CSCM is cooled down to 650 °C (SESMR condition), making the thermal power \( P_{\text{th7}} \) available. It is worth stressing that the adopted scheme may represent either a dual fluidized bed reactor with solid recirculation or a
packed bed configuration. From an energy perspective, CSCM circulation can explain the thermal power required by the calcination (3), and the cooling of the regenerated CSCM makes $P_{th,6}$ available. Therefore, the difference is that in dual inter-circulating fluidized bed conditions, this representation coincides with the physics of the process, whereas, in the case of packed bed reactors, the flow charts reproduce only the thermal power exchange, as the solid is not actually circulated.

(h) The absorbed CO$_2$ is released once the CSCM regeneration is complete. To store CO$_2$, it should be refrigerated and compressed. For the sake of simplicity, in this phase, only the refrigeration stage is considered to preliminarily assess the thermal power made available, $P_{th,8}$.

(i) Depending on the application, the fuel (H$_2$, CH$_4$, H$_2$O, CO, or CO$_2$) can be subjected to a certain degree of cooling, making the thermal power $P_{th,9}$ available.

Figure 1. Scheme of the energetic model of SESMR.

2.3. Power Contribution Assessment through the Energetic Model of SESMR

To perform an energy assessment of the experimented system, the thermal powers—required or made available—were quantified via the calculation of the thermodynamic conditions of the different involved substances (Appendix A). Thus, the different power
contributions were evaluated. The first is the power $P_{th,1}$, needed at Step (a), which can be determined by Equation (4), as follows:

$$P_{th,1} = \dot{m}_a \left( h_{a, out} - h_{a, in} \right)$$  \hspace{1cm} (4)

where $\dot{m}_a$ is the water mass flow rate, while $h_{a, out}$ and $h_{a, in}$ are the enthalpy of the steam at the evaporator outlet and water at the inlet, respectively.

Subsequently, the state of the mixture at Step (b) was determined as follows:

1. The mass conservation was applied, as given by Equation (5), considering the species (steam and methane) involved in the mixture:

$$\sum v_j M_j = 0$$  \hspace{1cm} (5)

where $v_j$ are stoichiometric coefficients (positive if $j$ is a product, negative if $j$ is a reactant) and $M_j$ is the molecular mass of species $j$.

2. The thermal power needed by the methane–steam mixture to reach the state required to enter the SESMR reactor is given by Equation (6), as follows:

$$P_{th,2} = \int_{T_{c, in}}^{T_{c, out}} \sum F_j c_{p,j}(T)dT$$  \hspace{1cm} (6)

where $F_j$ is the molar flow rate of the $j$ substance of the mixture and $c_{p,j}(T)$ heat at a constant pressure of species $j$ ($c_{p,j}$) as a function of absolute temperature ($T$).

3. The thermal power $P_{th,3}$ is associated with the overall reaction of SMR (1) and WGS (2). Thus, it is given by Equation (7):

$$P_{th,3} (T) = \sum F_j \Delta H_f^{\circ \text{SMR}, 298}$$  \hspace{1cm} (7)

where $\Delta H_f^{\circ \text{SMR}, 298}$ is the heat of formation of chemical species involved in both SMR and WGS reactions. $P_{th,3}$ represents the net difference between the power needed by SMR (highly endothermic) and that provided by WGS (slightly exothermic). The energy balance return that the energy requires by the reforming is higher than that made available by the WGS; thus, $P_{th,3}$ (8) expresses a power contribution required by the system. The fuel at the inlet is constituted by a mixture of steam and methane, while that at the outlet of the reactor is a reformed gas composed of $\text{H}_2$, CO, $\text{CO}_2$, $\text{CH}_4$, and $\text{H}_2\text{O}$.

4. $P_{th,4}$ (8) represents the thermal power provided by the carbonation which is an exothermic reaction. This energy contribution can be evaluated as follows:

$$P_{th,4} (T) = F_{\text{CO}_2} \Delta H_f^{\circ \text{CBN}, 298}$$  \hspace{1cm} (8)

where $F_{\text{CO}_2}$ is the absorbed $\text{CO}_2$ molar flow rate and $\Delta H_f^{\circ \text{CBN}, 298}$ is the heat of reaction associated with the carbonation process.

5. As above reported, the sorbent needs the thermal power $P_{th,5}$ to reach the condition required by the sorbent regeneration phase, as given by Equation (9).

$$P_{th,5} (T) = \int_{T_{c, in}}^{T_{c, out}} \sum F_{\text{CaCO}_3} \frac{\dot{m}_{\text{CaCO}_3}}{x_{\text{CaCO}_3}} c_{p,CSCM}(T)dT$$  \hspace{1cm} (9)

where $x_{\text{CaCO}_3}$ is the $\text{CaCO}_3$ molar fraction in combined sorption–catalyst material (CSCM) and $c_{p,CSCM}(T)$ is the specific heat at a constant pressure of the CSCM.
6. The thermal power required by the calcination $P_{th,6}$ is given by Equation (10):

$$P_{th,6}(T) = F_{CO_2} \left(-\Delta H^\circ_{f,CBN \ 298}\right)$$  \hspace{1cm} (10)

where $\left(-\Delta H^\circ_{f,CBN \ 298}\right)$ is the heat of formation related to the calcination stage.

7. At the end of the calcination stage, the CSCM reaches a temperature equal to 950 °C ($T_{g,in}$) in case of severe calcination. Then, CSCM should be cooled down to 650 °C ($T_{g,out}$) in such a way that the reforming stage takes place again. Thus, during Step (f), the CSCM cooling makes the thermal power $P_{th,7}$ available, as given by Equation (11):

$$P_{th,7}(T) = \int_{T_{g,in}}^{T_{g,out}} F_{CO_2} \cdot \frac{\rho_{CSCM}(T)}{x_{CaO}} \cdot dT$$  \hspace{1cm} (11)

where $x_{CaO}$ is the CaO molar fraction of CSCM.

It is important to notice that $P_{th,3}, P_{th,4}, P_{th,5}, P_{th,6}$ and $P_{th,7}$ take place simultaneously during the SESMR processes. The disaggregation was performed to evaluate the thermal power requirements and availability for an energy assessment.

8. Once the CO$_2$ has been released by the sorbent (g) it must be cooled from $T_{h,in}$ (950 °C) to $T_{h,out}$ and compressed to complete the storage stage. The analysis performed here was pushed until the step that precedes the intercooled compression as the main scope was to define the main source of thermal power needed and made available by the system. In particular, the CO$_2$ cooling provides $P_{th,8}$ to the surrounding environment, as in Equation (12):

$$P_{th,8}(T) = \int_{T_{h,in}}^{T_{h,out}} F_{CO_2} \cdot c_{p,CO_2}(T) \cdot dT$$  \hspace{1cm} (12)

9. In addition, once the fuel has left the reactor, it should be cooled from $T_{i,in}$ (650 °C) to $T_{i,out}$ (ambient temperature) to make its condition suitable for storing the hydrogen. The power $P_{th,9}$, made available by this cooling, is given by Equation (13):

$$P_{th,9}(T) = \int_{T_{i,in}}^{T_{i,out}} \sum F_j \cdot c_{p,j}(T) \cdot dT$$  \hspace{1cm} (13)

Once the power contributions were evaluated, three performance criteria were developed to assess the goodness of the process from an energy point of view.

The ratio between H$_2$ and CH$_4$ chemical energy is reported by $\xi_{H_2,A}$ in Equation (14):

$$\xi_{H_2,A} = \frac{m_{H_2} H_{iH_2}}{m_{CH_4} H_{iCH_4}}$$  \hspace{1cm} (14)

being representative of the direct recovery of thermal energy through the reforming process [72].

A further indicator of the energy performance is $\eta_{H_2,B}$ in Equation (15), which also considers the power required by the process $P_{th,req}$ in Equation (16):

$$\eta_{H_2,B} = \frac{m_{H_2} H_{iH_2}}{m_{CH_4} H_{iCH_4} + P_{th,req}}$$  \hspace{1cm} (15)

$$P_{th,req} = \frac{(P_{th,1} + P_{th,2} + P_{th,3} + P_{th,5} + P_{th,6})}{\eta_b}$$  \hspace{1cm} (16)
where $\eta_b$ is the burning efficiency of the generator adopted to obtain the required thermal power.

The total recovered power was calculated by Equation (17), as follows:

$$P_{\text{th,rec}} = (P_{\text{th,4}} + P_{\text{th,7}} + P_{\text{th,8}} + P_{\text{th,9}}) \eta_{\text{rec}}$$

(17)

where $\eta_{\text{rec}}$ is the efficiency of the recovery system. The third indicator $\eta_{H_2,C}$ can be obtained by Equation (18), as follows:

$$\eta_{H_2,C} = \frac{m_{H_2} H_i}{m_{CH_4} H_i + P_{\text{th,req}} - P_{\text{th,rec}}}$$

(18)

### 3. Results

The developed model allows for deeper insight into the energy characterization of the process, thus assessing the thermal power requirements and availability in a SESMR/sorbent-regeneration process. This is the first step to carry out a future optimization of the SESMR process, which is based on the highly energy-intensive SMR.

Due to the large experimental data set, the model was used to provide a detailed energy assessment of the SESMR/sorbent-regeneration process. In particular, the first model input concerned the molar flow rates of the involved substances. A scale factor was used in the energetic model, which multiplies the actual molar flow rates involved in the bench-scale experiments (see Section 2.1). As a matter of fact, the molar flows were rather low for an energy evaluation. The adopted scale factor allowed for referring the whole integrated system to a fed CH$_4$ mass flow rate equaling 1 g/s. In this regard, even though N$_2$ was fed as a carrier gas together with CH$_4$ and steam in the actual SESMR/sorbent regeneration experimental tests [70,72], the presence of N$_2$ was not considered in the model and was a carrier gas added to the gaseous mixture only for bench-scale measurability purposes.

The first evaluation regarding CaO15Ni10 performances over long reforming steps (see Section 2.1) was carried out by a comparison between the first SESMR session and the following processes (SMR+WGS, devoid of any sorption-enhancing effect). Despite the methane conversion of both SESMR and SMR+WGS leading to similar results (Figure 2a), the advantages introduced by CO$_2$ sorption emerged by observing the hydrogen molar fraction in the fuel. Indeed, Figure 2b shows that the SESMR allowed for achieving a higher H$_2$ molar concentration on a dry, dilution-free basis in comparison to the SMR+WGS. When SESMR took place, an average H$_2$ molar fraction of about 90 mol% was observed, which is considerably higher than that obtained through SMR+WGS (about 77 mol%). The reported H$_2$ molar fractions correspond to about 70 mol% and 55 mol%, respectively, on a wet, dilution-free basis (Figure 2b). The energetic model can be considered to have two positive benefits of SESMR towards H$_2$ concentration since the sorption-enhancing process reduces the CO$_2$ concentration in the obtained fuel gas and simultaneously drives reactions to further H$_2$ production.

The advantages introduced by SESMR were also clear from an energy point of view. Indeed, in Figure 3, the trend of $\xi_{H_2,A}$ can be observed, which expresses the hydrogen chemical energy at the SESMR exit versus the methane one at its inlet (14). Therefore, Figure 3 shows that the SESMR allowed for achieving higher values in comparison to the one obtained with SMR+WGS. In particular, with SESMR, the resulting average value is around 115%, whereas, for SMR+WGS, the trend oscillated more, with $\xi_{H_2,A}$ ranging between 93% and 103%. It is worth noting that in the SESMR situation, some cases (between cycles 20 and 45) were relatively far from the average trend. This behavior correlated with a slight negative deviation from the set point of the CH$_4$ inlet flow rate, which experimentally occurred during the long reforming steps between cycles 20 and 45. This led to obtain less CH$_4$ and H$_2$ in the gas produced during the affected cycles in comparison to regular operations, and, therefore, to a slight overestimation of CH$_4$ conversion (Figure 2a). In addition, a slight decrease of the H$_2$ fraction on a wet basis was even amplified on a relative basis (Figure 2b).
Once the $\xi_{H_2,A}$, was evaluated, the next step for a more thorough energy performance assessment was the characterization of various thermal power contributions required by the process $P_{th,req}$ (16). Figure 4 reports the percentage distribution of thermal power required by different stages.
The knowledge about the total power required by the system $P_{th,req}$ allowed for evaluating the performance index $\eta_{H_2,B}$ (Figure 5a), which represents the energy efficiency of the integrated process while taking into account the power required by different stages of the system. Figure 5a shows $\eta_{H_2,B}$ (15) for the SESMR and SMR+WGS cases. It can be observed that SESMR ensured higher values in comparison to those obtained by SMR+WGS. Even though the differences ranged by a few percentage points, the separation of CO$_2$ and H$_2$ (almost pure streams) could not be neglected. The small difference of $\eta_{H_2,B}$ between the SESMR and SMR+WGS cases was ascribable to the fact that higher $P_{th,req}$ resulted in SESMR than in SMR+WGS due to the regeneration of the sorbent. Nevertheless, as can be observed in Figure 1 in the SESMR process, several stages made thermal power $P_{th,rec}$ (18) available, and that can reduce the total energy required by the system. These contributions were related to the following factors:

1. Carbonation;
2. Thermal power made available by CSCM cooling;
3. Thermal power recovered through CO$_2$ cooling;
4. Thermal power recovered through reformed fuel cooling.

On the contrary, for SMR+WGS, there were no stages for recovering thermal power, except for the thermal power recoverable through the fuel cooling. Thus, also considering the thermal power recovered, the performance index $\eta_{H_2,C}$ (17) was evaluated (Figure 5b). It represents the energy efficiency of the integrated process while taking into account the power contributions required and recoverable in the different stages of the integrated system.
It can be seen that the recovered power allowed for reducing the impact of the power required by the system on the energy efficiency, leading to a $\eta_{\text{H}_2,C}$ of SESMR equal to an average value of 79%, with a remarkable increase in comparison to $\eta_{\text{H}_2,B}$. Thus, SEMSR allowed for obtaining $\eta_{\text{H}_2,C}$ values significantly higher than those of SMR+WGS (72%), demonstrating the effectiveness of sorption-enhancing, not only in terms of avoided CO$_2$ but also from an energy point of view. In fact, it should be emphasized that the best energy performance ensured with the SESMR process was accompanied by a drastic reduction in CO$_2$ emissions.

The effective integration between the available thermal power and that required for the SESMR process can be seen in Figure 6, where the temperature trend is shown for the different stages as a function of associated thermal power. Figure 6 shows that the thermal power requirements (red line) at different stages can be covered considerably by the thermal power provided by other stages:

1. The carbonation makes $\text{P}_{\text{th},4}$ available, which reduces the thermal power needed by the reforming process ($\text{P}_{\text{th},3}$).
2. The thermal power $\text{P}_{\text{th},7}$, provided by CSCM cooling, can be used to cover part of the remaining $\text{P}_{\text{th},3}$ and the final stage of the fuel heating up prior to entering the reactor ($\text{P}_{\text{th},2}$). This means that the thermal power needed to heat up the sorbent for its regeneration can be partially recovered.
3. The cooling of reformed fuel exiting the reactor ($\text{P}_{\text{th},8}$) provides thermal power needed by the first part of the fuel heating up ($\text{P}_{\text{th},2}$) and by the steam superheating.
4. The CO$_2$ cooling ($\text{P}_{\text{th},9}$) partially covers the water vaporization stage. A further contribution can be provided by $\text{P}_{\text{th},9}$.

Therefore, the water economization, the main part of vaporization (part of $\text{P}_{\text{th},1}$), and the calcination ($\text{P}_{\text{th},6}$) emerged as the only stages that would remain uncovered. It is worth to stress that the trends reported in Figure 6 did not consider the heat exchange losses (which could be included in a future improved model) or evaluate the thermal power recovered from reformed fuel and captured CO$_2$ cooling, fixing their minimum temperature at 150 °C. Indeed, the main scope of the study was to perform a preliminary energy characterization of the bench-scale system (1 g/s of fed CH$_4$) to open the way for the optimization of possible industrial SESMR applications.

Figure 5. Hydrogen production efficiency (a) $\eta_{\text{H}_2,B}$ without thermal energy recovery and (b) $\eta_{\text{H}_2,C}$ with thermal energy recovery for SESMR and SMR.
Figure 6. Temperature as a function of thermal power for different SESMR stages (red lines are the required powers, blue lines are the available powers).

4. Conclusions

In the present study, a comprehensive thermodynamic model was developed for the stages constituting sorption enhanced steam methane reforming (SESMR) systems in order to assess their energy performance. The model was applied to an actual SESMR bench-scale plant, scaling up involved flow rates to get proper quantities for an energy evaluation. The experimental information provided by previous campaigns allowed for carrying out a detailed energy characterization of the system. Moreover, the benefits introduced by the SESMR were assessed, considering reforming stages in which the sorbent was saturated (and, consequently, assimilable to steam methane reforming and water gas shift (SMR+WGS) without sorption-enhancing). In particular, the following results were observed:

1. Although the methane conversions were quite close, SESMR presented a higher hydrogen molar fraction on a dry (90 mol%) and wet basis (70 mol%), in comparison to SMR+WGS without CO$_2$ adsorption, in agreement with the experimental results.
2. SESMR allowed for achieving a higher value of the first performance index $\xi_{H_2,A}$, defined as the ratio between hydrogen and methane chemical energy. The results show that SESMR presented an average $\xi_{H_2,A}$ equal to 115%, while that of SMR fluctuated between 105% and 95%;
3. Considering the power required by the system ($\eta_{H_2,B}$), it was observed that even though the advantages offered by SESMR were still present, they were limited to a few percentage points. This was mainly due to the higher thermal power required by the SESMR loop for sorbent regeneration.
4. The final step of the study demonstrated that in SESMR, a high fraction of the required thermal power could be recovered from energy contributions provided by the carbonation process, sorbent, CO$_2$, and fuel cooling. This allowed for reducing the power demand of the system and improving the plant energy performance. Indeed,
considering the contributions of recovered thermal power in the energy performance \( \eta_{H_2,C} \), an average value of 79% was observed for SESMR, while for SMR+WGS a maximum value of 72% was observed.

Therefore, the proposed model allows for performing a comprehensive energy analysis of the system, demonstrating that SESMR has benefits not only in terms of hydrogen production and carbon dioxide separation but also referring to the energy performance of the integrated system. Indeed, despite the higher thermal power required in comparison to the SMR+WGS process at the same process conditions (650 °C, 1 atm), the possibility to recover a large fraction of this power improves the energy performance of the entire SESMR/sorbent-regeneration process.

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**Nomenclature**

**Acronyms**

| Acronym | Description |
|---------|-------------|
| CBN     | Carbonation |
| CSCM    | Combined catalyst sorption material |
| SESMR   | Sorption-enhanced steam–methane reforming |
| SMR     | Steam–methane reforming |
| WGS     | Water–gas Shift |

**Symbols**

| Symbol | Description |
|--------|-------------|
| \( c_{p,j} \) | Specific heat at constant pressure of species \( j \) [kJ/(kmolK)]/[kJ/kgK] |
| \( F \) | Molar flow rate of species \( j \) [kmol/s] |
| \( h \) | Specific enthalpy [kJ/kg] |
| \( M \) | Molecular masses [kg/kmol] |
| \( \dot{m} \) | Mass flow rate [kg/s] |
| \( P_{th} \) | Thermal power [W] |
| \( T \) | Temperature [K] |
Subscripts
(a) Vaporization phase
(b) Steam/methane mixing phase
(c) Fuel heating phase
(d) Sorption-enhanced reforming phase
(e) Sorbent heat up for regeneration
(f) Calcination phase
(g) Sorbent cooling phase
(h) CO\(_2\) cooling phase
(i) Fuel cooling phase
\(j\) Generic species
\(\text{in}\) Inlet
\(\text{out}\) Outlet

Greek symbols
\(\Delta H^\circ_f, i\) Standard heat of formation of \(i\)-substance [kJ/kmol]
\(\eta_{H_2,B}\) \(H_2\) production efficiency (without energy recovery)
\(\eta_{H_2,C}\) \(H_2\) production efficiency (with energy recovery)
\(\nu\) Stochiometric coefficient [kmol]
\(\xi_{H_2,A}\) \(H_2\) vs. \(CH_4\) chemical energy

Appendix A

This appendix summarizes the calculation methods for thermodynamic properties considered in this work.

Specific heat at a constant pressure of species \(j\) \((c_p,j)\) as a function of absolute temperature \((T)\) was calculated by (A1), with constants for each species \((a_j, b_j, c_j, d_j)\), as reported in Table A1.

\[
c_{p,j}(T) = a_j + b_j T + c_j T^2 + \frac{d_j}{T^2} \quad \text{with } c_{p,j} \text{ in } \frac{\text{cal}}{\text{mol K}} \text{ and } T \text{ in K}
\]  

(A1)

Table A1. Constants for \(c_{p,j}\) calculation as a function of absolute temperature, according to A1 (phases: (c) = crystalline; (g) = gas).

| \(j\)    | Phase | \(a_j\) (cal mol\(^{-1}\) K\(^{-1}\)) | \(b_j\) (cal mol\(^{-1}\) K\(^{-2}\)) | \(c_j\) (cal mol\(^{-1}\) K\(^{-3}\)) | \(d_j\) (cal mol\(^{-1}\) K\(^{3}\)) | Validity (K) | Ref. |
|----------|-------|--------------------------------------|--------------------------------------|--------------------------------------|--------------------------------------|--------------|------|
| CaCO\(_3\) | (c)    | 19.68                                | 0.01189                              | 0                                     | \(-307,600\)                         | 273–1033     | [59] |
| CaO      | (c)    | 10.00                                | 0.00484                              | 0                                     | \(-108,000\)                         | 273–1173     | [59] |
| CO\(_2\) | (g)    | 10.34                                | 0.00274                              | 0                                     | \(-195,500\)                         | 273–1200     | [59] |
| CH\(_4\) | (g)    | 5.34                                 | 0.01150                              | 0                                     | \(-130,300\)                         | 273–1373     | [59] |
| CO       | (g)    | 6.60                                 | 0.00120                              | 0                                     | \(0\)                                | 273–2500     | [59] |
| H\(_2\)  | (g)    | 6.62                                 | 0.00081                              | 0                                     | \(0\)                                | 273–2500     | [59] |
| H\(_2\)\(_2\) | (g) | 8.22                                 | 0.00015                              | \(0.00000134\)                        | \(0\)                                | 300–2500     | [59] |
| Ni       | (c\(\alpha\)) | 4.26                                | 0.0064                               | 0                                     | \(0\)                                | 273–626      | [59] |
| Ni       | (c\(\beta\)) | 6.99                                | 0.00905                              | 0                                     | \(0\)                                | 626–1725     | [59] |
| NiO      | (c)    | 11.3                                 | 0.00215                              | 0                                     | \(0\)                                | 273–1273     | [59] |
| O\(_2\)  | (g)    | 8.27                                 | 0.000258                             | 0                                     | \(-187,700\)                         | 300–5000     | [59] |
| Ca\(_{12}\)Al\(_{14}\)O\(_{33}\) | (c\(\alpha\)) | 301.96                              | 0.0655                               | \(0\)                                | \(-5,530,000\)                      | 298–1300     | [60] |
| Ca\(_{12}\)Al\(_{14}\)O\(_{33}\) | (c\(\beta\)) | 228.52                              | 0.09844                              | \(0\)                                | \(0\)                                | 1300–1700    | [60] |

Concerning the CSCM, its specific heat \((c_{p,CSCM})\) as a function of \(T\) was calculated as the weighted average of \(c_{p,j}\) of its chemical components ((A2), Table A1). CSCM compositions (molar fractions \(x_j\)) depend on their chemical state (reduced or oxidized catalytic phase, calcinated or carbonated sorbent phase).

\[
c_{p,CSCM}(T) = \sum x_j c_{p,j}(T) \quad \text{with } j = \text{Ni}, \text{NiO}, \text{CaO}, \text{CaCO}_3, \text{Ca}_{12}\text{Al}_{14}\text{O}_{33}
\]  

(A2)

Let us consider the generic chemical reaction in (A3), where \(v_j\) is the stochiometric coefficient (positive if \(j\) is a product, negative if \(j\) is a reactant) and \(M_j\) is the molecular mass...
of species \( j \). The heat of that reaction as a function of temperature \((\Delta H_R(T))\) was calculated by Kirchoff (A4), with the standard heat of \((\Delta H^\circ_R)\) determined according to Hess’s law (A5), where \(\Delta H^\circ_{f,j}\) is the standard heat of formation of \( j \) shown in Table A2.

\[
\sum v_j M_j = 0 \quad \text{(A3)}
\]

\[
\Delta H_R(T) = \Delta H^\circ_R + \int_{298.15}^{T} \sum v_j c_{p,j}(T) dT \quad \text{(A4)}
\]

\[
\Delta H^\circ_R = \sum v_j \Delta H^\circ_{f,j}\quad \text{(A5)}
\]

| \( j \) | Phase | \( \Delta H^\circ_{f,j}\) \( (\text{kcal mol}^{-1} \text{ K}^{-1}) \) | Ref. |
|------|------|-----------------|------|
| CaCO\(_3\) | (c) | \(-289.5\) | [59] |
| CaO | (c) | \(-151.7\) | [59] |
| CO\(_2\) | (g) | \(-94.052\) | [59] |
| CH\(_4\) | (g) | \(-17.81\) | [59] |
| CO | (g) | \(-26.416\) | [59] |
| H\(_2\) | (g) | \(0\) | [59] |
| H\(_2\)O | (g) | \(-57.7979\) | [59] |
| Ni | (c\(\alpha\)) | \(0\) | [59] |
| Ni | (c\(\beta\)) | \(0\) | [59] |
| NiO | (c) | \(-58.4\) | [59] |
| O\(_2\) | (g) | \(0\) | [59] |

Table A2. Standard heats of formation \(\Delta H^\circ_{f,j}\) in (A4) (phases: (c) = crystalline; (g) = gas).

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