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Electrified CO₂ valorization driven by direct Joule heating of catalytic cellular substrates

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ABSTRACT

The growing environmental concerns have driven catalytic CO₂ valorization as a forward-looking solution to mitigate the carbon footprint of valuable chemical products. CO₂ conversion processes into synthesis gas, such as CO₂ reforming of methane (CRM) or reverse water-gas shift (RWGS), may have a strategic role for the future sustainable production of chemicals and energy carriers. However, fuel combustion to supply the heat of the associated endothermic reactions would result in unwanted CO2 emissions, frustrating the overall objective. Electrification of the endothermic processes may represent the technological solution to such an issue. Here we report a promising approach for the direct electrification of the CO₂ reforming of methane (eCRM) and reverse water-gas shift (eRWGS) processes in washcoated structured reactors. We employ catalytically activated opencell foams that provide optimal heat and mass transfer properties and serve as Joule heating substrates for the catalytic conversion of CO2 via reaction with methane or hydrogen. The proposed reactor system with Jouleheated Rh/Al₂O₃-coated foam exhibited excellent catalytic and electrical stability for more than 75 h, operating up to 800 °C and approaching equilibrium conversion at high space velocity, i.e., GHSV of 600 and 100 kNl/kgcat/h for eRWGS and eCRM, respectively. Such a reactor concept has potential to ensure remarkably low specific energy demand for CO₂ valorization. Assuming an optimized process configuration approx. 0.7 kWh/ Nm³_{CO2} is calculated for eRWGS. By replacing fuel combustion with Joule heating driven by renewable electricity, the electrified CO₂ valorization processes provide an important approach for dealing with the intermittent nature of renewable sources by storing the energy in chemicals with a low carbon footprint.

1. Introduction

To meet the demand of a growing global population, unprecedented amounts of fossil fuels are used as a source of energy and feedstocks to produce necessary chemicals, releasing significant amounts of greenhouse gas and other pollutants. It is estimated that the energy consumption by the chemical industry reached 43 EJ/y in 2015, together with roughly 3.3 Gt CO₂eq emissions released to the environment, which however shows a constantly increasing trend [1]. As a result of the growing environmental concerns, CO₂ valorization has been proposed as a promising path to mitigate the carbon footprint of valuable chemical products (e-fuels, e-polymers, e-chemicals). In this context, the CO₂ reforming of methane (CRM, Eq. (1)) and the reverse water–gas shift (RWGS, Eq. (2)) reactions are interesting options for syngas production from captured CO₂ [2–4].

$$CRM: CO_2 + CH_4 \rightleftharpoons 2CO + 2H_2 \quad \Delta H_r^o = 247 \text{ kJ/mol}$$
(1)

 $RWGS: CO_2 + H_2 \rightleftharpoons CO + H_2O \qquad \Delta H_r^o = 41 \text{ kJ/mol}$ (2)

However, the endothermic nature of both reactions indicates that thermal energy is required to sustain them, which must be provided at high temperature to allow optimal process performances. In industrial-scale applications, the heat of reaction is typically supplied by fuel combustion: for CO_2 valorization processes this is paradoxical, negatively affecting the overall CO_2 balance. It is estimated that about half of the CO_2 emissions in the chemical industry are related to fuel combustion for heat supply [5].

To effectively mitigate the carbon footprint of CO_2 conversion routes, the primary energy source needs to be changed [6,7]. The electrification of catalytic processes is a promising concept that offers great potential to utilize excess renewable energy and mitigate CO_2 emissions [8–11].

In the context of CO_2 valorization processes, electrochemical reduction of CO_2 (CO_2R) enables the direct utilization of electric energy

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into CO₂ conversion, however, it is important to note that the solid oxide electrolyzer, i.e. the only technology approaching commercialization with over one year durability demonstrated [12], is characterized by an overall specific energy demand of 6–8 kWh/ Nm_{CO2}^3 [13]. On the other hand, power-to-heat approaches, such as induction heating, microwave heating and Joule heating, enable direct thermal energy supply without changing the reaction mechanism [14–16]. In principle, an intrinsically higher thermal efficiency (theoretically 100% [17]) can be expected from Joule heating as it is the only one that enables the direct transformation of electricity into thermal energy. Moreover, Joule heating is also characterized by fast dynamics and it has been proven that heat can be delivered at significantly high temperatures [18-20]. Process environmental and economic calculations suggest that Joule-heated CO2 utilization processes based on renewable electricity are promising to reduce the carbon footprint of chemical products in the near future [21.22].

From the reactor engineering standpoint, one key issue is how to implement Joule heating in the design of electrified catalytic reactor. It is possible to directly heat up conductive catalyst pellets by Joule effect [23,24], however it is practically difficult to control the size of the contact area between pellets to enable electrical continuity [25]. On the other hand, the Joule heating concept is well applicable to structured catalysts, whose continuous solid matrix can ensure electrical continuity. For example, it is feasible to adopt catalytically activated commercial heating resistances with different shapes located inside the reactor [26–28], or integrate the electrical resistances (heating wires) within the channels or the framework of ceramic honeycomb monolith catalysts [29,30]. Joule heating can also be directly applied to the reactor tube, washcoated with catalyst on the inner wall [8,31]. Such configurations ensure excellent heat transfer, since the heat source is directly in contact with the catalyst; however, they are associated with significant external mass transfer limitations [32]. As an alternative solution, Joule-heated open-cell foams, characterized by high specific surface areas and excellent transport properties, have potential to overcome heat and mass transfer limitations in the electrified reactors [33-37].

These considerations motivated us to explore the electrified CO_2 reforming of methane (eCRM) and the electrified reverse water–gas shift (eRWGS) reactions by direct Joule heating of catalytically-activated open-cell SiC foams. In the present work, the Joule heating performance, catalytic performance, energy performance and CO_2 valorization performance of the electrified reactor were assessed for the eCRM and eRWGS processes.

2. Materials and methods

As in previous works from our group [33,37], the adopted SiSiC foam ($d_{foam} = 3.2$ cm, Erbicol, CH) has a cell diameter (d_{cell}) of 3.32 mm and a strut diameter (d_{strut}) of 0.61 mm, a total porosity estimated by ethanol picnometry of 0.88, and a surface to volume ratio evaluated on the bare geometry (Sv) of 740 m⁻¹ [33]. The bulk foam material was characterized by X-ray diffraction: the analysis showed that both Si and SiC phases are present. In this work, SiSiC foams with length of 9.9 cm and 8 cm were utilized.

Rh-based catalysts are reported to be active for both reforming and reverse water-shift reactions, and have also demonstrated remarkable ability to resist coking even in demanding conditions such as CO_2 reforming of methane. [38–41]. In order to activate the SiSiC foam for catalytic reactions, a 1% Rh/Al₂O₃ catalyst was prepared in a first step via a wet-impregnation method using γ -Al₂O₃ powder (Sasol, PUR-ALOX) as morphological support and rhodium (III) nitrate solution (Rh 10–15% w/w, Alfa Aesar) as rhodium precursor [33,37]. Afterwards, a washcoating slurry was prepared by mixing the catalyst powders with PVA (polyvinyl alcohol, Sigma-Aldrich), glycerol (Sigma-Aldrich) and deionized water via magnetic stirring and ball-milling (50 rpm) for 24 h. The SiSiC foam washcoating was achieved by dipping the foam in the catalyst slurry, followed by spinning (1000 rpm for 10 s) to remove the

excess material and by flash drying in oven at 350 °C for 5 min. More details on the catalyst/slurry preparation and foam washcoating procedures can be found in our previous works [33,37]. The obtained catalytic activated SiSiC foams are shown in SI (Section S1, Fig. S1 and Fig. S2).

As shown in Fig. 1, the washcoated SiSiC foam was placed inside a tubular stainless-steel reactor (OD = 5 cm), as previously described [33,37]. The SiSiC foam was connected through electric contactors to an AC rheostat ($V_{max} = 50 \text{ V}$, $I_{max} = 100 \text{ A}$, Belotti Variatori) hooked to the electric grid (230 V, 50 Hz). The rheostat allowed to vary the voltage and measure the current intensity. A ceramic tube (ID = 3.5 cm) was inserted between the foam and the stainless-steel reactor tube to insulate the system. To connect the foam with the AC power generator, home-made electric contactors were adopted, these elements are made of stainlesssteel cylindrical plates, provided with some holes for gas flow and welded to a 1/4' stainless steel pipe. A thin layer of copper foam (Alantum, pore size 800 µm, 1 mm thickness) was placed between the SiSiC foam and the electrical contactors to minimize contact resistances. Moreover, a more robust external metallic frame structure compared to previous works [33,37] was adopted to apply additional force to the electrical contactors in order to compensate the effect of their thermal expansion. K-type thermocouples, electrically insulated by ceramic thermowells (dense alumina, $d_{out} = 3 \text{ mm}$, $d_{in} = 2 \text{ mm}$), were placed inside the upstream and downstream electric contactors to measure the temperatures at the upper and bottom cross sections of the foam, the measured temperatures were noted as Tup and Tdown, respectively. Tcap was measured by a fine thermocouple placed in a quartz capillary (inner diameter of 1 mm, outer diameter 2 mm) and located at 10 mm from the centerline along the radial coordinate, through the Cu felt and in direct contact with the bottom of the foam. Another sliding thermocouple (T_{slide}) was placed inside a thermowell welded at the outer reactor wall. Temperatures were measured with a resolution of 1 cm, with a total length of 15 cm equal to the reactor length.

The electrified CO_2 reforming of methane and reverse water–gas shift runs were performed at ambient pressure, at different temperatures, space velocity and feed composition. A complete list of the experimental runs is reported in SI (Section S3, Table S1 and Table S2). The gases were fed to the system individually by means of mass flow controllers (Brooks 5851). Catalytic performance at different T_{cap} temperatures was



Fig. 1. Schematic representation of the electrified reactor layout. The red dots indicate where the temperatures are measured. The illustration is not to scale.

evaluated by adjusting the input voltage, the electric current selfadjusting according to the resistance of the system. The outlet gas composition was analyzed using an online micro-GC (Agilent, 900 Micro GC) after removing water from the products by a condenser. An inert gas (nitrogen) was directly fed to the micro-GC through a by-pass as an internal standard. A satisfactory closure of the carbon balance (\pm 5%), considering CH₄, CO and CO₂ species, was obtained in all the tests. Water in the outlet stream was quantified according to the extents of reaction calculated by considering the other measured outlet species.

3. Results and discussion

3.1. Joule-heating of bare SiSiC foam

Before conducting Joule-heated catalytic experiments using catalyst washcoated SiSiC foams in a stainless-steel tube reactor, it is essential to evaluate the Joule heating performance of the bare foam. For this purpose, a quartz tube reactor was utilized to investigate the thermal behavior of the foam during the Joule heating process. Fig. 2 illustrates the Joule heating of a bare SiSiC foam loaded in a quartz tube, the foam was electrically connected through homemade contactors to an AC rheostat. To minimize contact resistances, a thin layer of copper foam was placed between the SiSiC foam and the electrical contactor. To measure the foam temperature, one thermocouple was placed inside the reactor with the foam. Another thermocouple was located outside the reactor wall.

Fig. 2 (a) shows the loaded foam at room temperature. Thanks to the interconnected geometry and suitable bulk resistivity of the SiSiC foam, it can be directly Joule heated to the relevant temperatures for electrified CO₂ reforming of methane and electrified reverse water–gas shift reactions. The adopted SiSiC foam ensures uniform power distribution and stable electrical resistance (approx. 0.3 ohm) at 700 °C in 25 Nl/h N₂ flow, as shown in Fig. 2(b).

3.2. Electrified CO₂ conversion: eCRM and eRWGS

Catalytic tests were performed by progressively increasing the electrical power input. Results in Fig. 3(a) show that the structured catalyst could be heated by the Joule effect up to the high temperatures relevant for CRM and RWGS reactions, thanks to the proper geometry and bulk resistivity of the SiSiC foam. The washcoated foam has an average catalyst layer thickness of 40 μ m and a catalyst density of 39.3 g/L considering that only 4/5 of the foam volume was washcoated (Fig. S1). An almost linear correlation was observed between the input power and



Fig. 2. Joule heating of the bare SiSiC open-cell foam ($d_{foam} = 3.2 \text{ cm}$, $L_{foam} = 9.9 \text{ cm}$, Erbicol, CH) in a quartz tube. (a) SiSiC foam at room temperature, the foam is connected to an AC power supplier through homemade electrical contactors. (b) Joule heated SiSiC foam with a uniform temperature distribution at 700 °C in 25 Nl/h N₂ flow, the gas flows downward from top to bottom.



Fig. 3. Thermal performance of the electrified reactor. (a) Measured T_{cap} temperature as a function of input power during electrified CO_2 reforming of methane (eCRM) and electrified reverse water–gas shift (eRWGS) runs. (b) Axial wall temperature profiles measured by T_{slide} during eCRM and eRWGS runs at the same T_{down} temperature of 700 °C. 1 %Rh/Al₂O₃ washcoated SiSiC foam ($L_{foam} = 9.9$ cm) with catalyst density of 39.3 g/L. eCRM: $CO_2/CH_4 = 4$, GHSV = 100 kNl/kg_{cat}/h, ambient pressure. eRWGS: $H_2/CO_2 = 2.25$, GHSV = 100 kNl/kg_{cat}/h, ambient pressure.

the measured T_{cap} temperature.

As illustrated in (Fig. 3(b)), the axial temperature profile of the reactor wall exhibited an increasing trend along the foam length, reaching a maximum in correspondence of the foam bottom section. Such a profile is consistent with the foam acting not only as the catalyst support but also as the Joule heating element [36]. A similar trend was observed at all the investigated temperatures (Fig. S3).

The measured CO₂ conversions closely approach the thermodynamic equilibrium calculated with T_{cap} in a wide range of conditions for both eCRM and eRWGS (Fig. 4). Such a close correspondence provided us confidence that T_{cap} is well representative of the outlet temperature from the catalytic foam. The obtained results in this work were compared to the literarure reports on eCRM and eRWGS with Joule heating in Table S3.

The outlet product distribution and H₂/CO ratio reported as a function of T_{cap} (Fig. 5) confirms the close approach to equilibrium above 675 °C. Noteworthy, at these high temperature conditions any CH₄ formation could hardly be detected during eRWGS runs, in line with thermodynamics. The good catalytic performance can be attributed to the following reasons: (i) the intimate contact between the catalyst phase and the heating element results in excellent heat transfer properties compared to traditional fuel combustion heating, where heat is generated outside of the reactor; (ii) the open-cell foam provides excellent mass transfer properties due to tortuous flow in its porous structure [42].

From entalphy balances it was found that the heat dissipation, i.e., the main source of power loss, varied almost linearly with the reactor temperature (T_{cap}) and was independent from the reaction chosen (Fig. 6(a)); it was thus ascribed to heat conduction through the thermal insulation layer of the reactor and natural convection. It is important to note that the ohmic losses in the wiring increase linearly with the input power. However, these losses were estimated to account for only approximately 3% of the total power input [36,37]. As for the reaction heat duty (Q, Eq. S8), more energy went into the reaction in the case of the eCRM process because of its strongly endothermic nature (Fig. 6(b)). This explains why eCRM requires a higher input power to reach the same temperature in comparison to eRWGS (Fig. 3(b)). As a result, the overall energy efficiency of the system, evaluated from the ratio of process heat duty to input power (Eq. S10), reached approx. 53% for the eCRM process, which was higher compared to 25% for the eRWGS process (Fig. 6(c)). In the SI, we predictively evaluated the performances of the two systems, showing an excellent agreement in terms of temperatures, conversions and thermal efficiencies (Section S6, Fig. S4, Fig. S5 and Fig. S6). Extensive research has demonstrated that utilizing direct Joule heating of structured catalysts within the reactor yields higher energy efficiency due to more selective heating of the catalysts [18,20,43-46],

(a)

Conversion [%]



Fig. 4. Catalytic performance of the electrified reactor. (a) Measured CO₂ conversion and methane conversion as a function of T_{cap} temperature during eCRM runs. (b) Measured CO₂ conversion and hydrogen conversion as a function of T_{cap} temperature during eRWGS runs. The experimental results (dots) are compared to the thermodynamic equilibrium conversions (lines). 1 % Rh/Al₂O₃ washcoated SiSiC foam (L_{foam} = 9.9 cm) with catalyst density of 39.3 g/L. eCRM: CO₂/CH₄ = 4, GHSV = 100 kNl/kg_{cat}/h, ambient pressure. eRWGS: H₂/CO₂ = 2.25, GHSV = 100 kNl/kg_{cat}/h, ambient pressure.

Fig. 5. (a) Outlet composition and (b) outlet H_2/CO ratio from electrified CO_2 reforming of methane (eCRM) runs. (c) Outlet composition and (d) outlet H_2/CO ratio from electrified reverse water–gas shift (eRWGS) runs. The experimental results (dots) are compared to thermodynamic equilibrium (lines). eCRM: $CO_2/CH_4 = 4$, GHSV = 100 000 Nl/kg_{cat}/h, ambient pressure. eRWGS: $H_2/CO_2 = 2.25$, GHSV = 100 000 Nl/kg_{cat}/h, ambient pressure.



Fig. 6. Energy performance of the electrified reactor. (a) The power loss of the system as a function of measured T_{cap} temperature during electrified CO₂ reforming of methane (eCRM) and electrified reverse water–gas shift (eRWGS) runs. (b) Reaction heat duty (Q) as a function of measured T_{cap} temperature during eCRM and eRWGS runs. (c) Energy efficiency (the ratio of reaction heat duty to input power) as a function of measured T_{cap} temperature during eCRM and eRWGS runs. 1 %Rh/Al₂O₃ washcoated SiSiC foam (L_{foam} = 9.9 cm) with catalyst density of 39.3 g/L. eCRM: CO₂/

850

 $CH_4 = 4, \ GHSV = 100 \ kNl/kg_{cat}/h, \ ambient \ pressure. \ eRWGS: H_2/CO_2 = 2.25, \ GHSV = 100 \ kNl/kg_{cat}/h, \ ambient \ pressure.$

in contrast to electric oven heating where heat is generated outside of the reactors. Further improvement of energy efficiency is still possible, for example, by increasing the total flow rate and by minimizing heat dissipations with better thermal insulation. More importantly, a significant improvement of the energy efficiency up to values close to 1 can be expected upon scale-up (Fig. S7), which facilitates quasi adiabatic operations [15,36].

The CO₂ valorization performances of the proposed foam-based

electrified reactor were evaluated based on the mole of reductant feed (CH₄ in eCRM and H₂ in eRWGS), catalyst and electrical energy inputs (Fig. 7). The eRWGS process reached a net CO₂ consumption (NC_{CO2}) of 1 mol_{CO2}/mol_{H2}, consistent with the reaction stoichiometry (CO₂/H₂ = 1, Eq. (2). In contrast, higher values of 1.3–1.6 mol_{CO2}/mol_{CH4} were achieved in the eCRM runs (Fig. 7(a)), thanks to the concurrent RWGS reaction where CO₂ (excess CO₂ as a result of CO₂ rich feed CO₂/CH₄ = 4) further reacted with the produced H₂ (Eq. (2)). The evidence for this aspect is provided by the K_p/K_{eq-rwgs} ratios computed during eCRM runs: as shown in Fig. S8, all such values were below 1, confirming that the RWGS reaction is indeed favoured in eCRM conditions.

The overall CO₂ conversion rate per catalyst mass (VR_{CO2}, Fig. 7(b)) reached 20 and 31 $\text{Nm}_{\text{CO2}}^2/\text{kg}_{\text{cat}}/\text{h}$ for eRWGS and eCRM, respectively, however, higher values could be expected with the increase of the feed CO₂/reductant ratio and especially the increase of the space velocity, since the system in all the investigated conditions was at thermodynamic equilibrium.

The eCRM process enables higher net CO₂ consumptions (Fig. 7(a)) and CO₂ conversion intensities (Fig. 7(b)), however, it requires greater electric energy input to reach the target reaction temperature in comparison to the eRWGS process (Fig. 3(b)). As a result, the specific energy demand for CO₂ valorization (SE_{CO2}, Eq. S14) fell in a similar range of 5.2–6.7 kWh/Nm³_{CO2} for both eCRM and eRWGS processes (Fig. 7(c)). The SE_{CO2} in eRWGS runs increased with input power as the heat loss played a major role at high temperatures; in contrast, a decreasing trend was observed in eCRM runs showing that a significant amount of energy was absorbed by the reaction.

The Joule-heated systems in principle enable significantly high energy efficiency [17]. Mathemathical model simulations demonstrated that approx. 95% efficiency can be reached in a 0.12 m diameter, 1.25 m long e-RWGS reactor operated at GHSV = 150 kNl/kg_{cat}/h with an input

power density of 6.5 MW/m³ (SI, Fig. S7), therefore based on this figure ($\eta^* = 0.95$) we calculate a quasi adiabatic specific energy (SE_{CO2}^* , Eq. S15) as low as 1.4 kWh/Nm³_{CO2}. Moreover, a big advantage could be gained by recovering the large fraction of sensible heat, i.e., by preheating the reactants, especially for eRWGS process ((Fig. 7(e)). In view of this, a second process integrated specific energy (SE_{CO2}^{**} , Eq. S18), evaluated by further considering the recover of 90% sensible heat [47], reached approx. 0.7 kWh/Nm³_{CO2} for eRWGS. In the eRWGS process, if the feed H₂ is sourced from water electrolysis (3.8 kWh/Nm³_{H2}) [48,49], it is possible to achieve an overall specific energy consumption of 4.5 kWh/Nm³_{CO2} for CO₂ valorization when operating with a recycle reactor. Such results are very attractive compared to solid oxide electrolyzers for CO₂ reduction (6–8 kWh/Nm³_{CO2}, [13]).

Since more energy is required to activate CH₄ than H₂ [3], the eCRM was observed to show higher SE_{CO2}^{**} values (2.3–2.6 kWh/Nm_{CO2}³) compared to eRWGS processes. In addition to CO₂ valorization, the eCRM process could be regarded as an environmental-friendly approach to activate methane. The decreasing trend of SE_{CO2}^{**} with temperature in eCRM runs could be explained by the concurrent RWGS reaction ((Fig. 7 (a)). This suggests that the SE_{CO2}^{**} is dependent on the feed gas composition and operating temperature, therefore further reduction of this value is in principle feasible, e.g., by feeding a more CO₂-rich gas mixture.

3.3. eRWGS: Effect of H_2/CO_2 feed ratio

As indicated by thermodynamics (*c.f.* equilibrium curves of Fig. 4), increasing the power input (and temperature) would have a positive, but limited effect on improving the CO_2 conversion during eRWGS runs. On the other hand, as previously discussed, the CO_2 conversion could be



Fig. 7. CO_2 valorization performance of the electrified reactor. (a) Net CO_2 consumption per mol of converted reductant (NC_{CO2}) as a function of input power during electrified CO_2 reforming of methane (eCRM) and electrified reverse water–gas shift (eRWGS) processes. (b) Overall CO_2 valorization rate (VR_{CO2}) as a function of input power during eCRM and eRWGS processes. (c) Specific energy consumption for CO_2 valorization (SE_{CO2}) as a function of input power during eCRM and eRWGS processes. (c) Specific energy consumption for CO_2 valorization (SE_{CO2}) as a function of input power during eCRM and eRWGS processes: experimental results based on input power (SE_{CO2}), the theoretical results considering an overall energy efficiency of 95% (SE_{CO2}^*), as well as a further revover of 90% sensible heat (SE_{CO2}^{**}). The energy distribution as a function of T_{cap} temperature during (d) eCRM and (e) eRWGS runs. 1 %Rh/Al₂O₃ washcoated SiSiC foam ($L_{foam} = 9.9$ cm) with catalyst density of 39.3 g/L. eCRM: $CO_2/CH_4 = 4$, GHSV = 100 kNl/kg_{cat}/h, ambient pressure. eRWGS: $H_2/CO_2 = 2.25$, GHSV = 100 kNl/kg_{cat}/h, ambient pressure.

further enhanced by increasing the H₂/CO₂ feed ratio. Our experimental results of CO₂ and H₂ conversions during eRWGS runs with different H₂/CO₂ feed ratios (T_{cap} = 800 °C, GHSV = 100 kNl/kg_{cat}/h, ambient pressure) are illustrated in Fig. 8. The experiments were performed using the same 1 %Rh/Al₂O₃ washcoated SiSiC foam (L_{foam} = 9.9 cm) with a catalyst density of 39.3 g/L. CO₂ conversion consistently increase with the H₂/CO₂ feed ratio up to 94.2%, which was achieved with an H₂/CO₂ feed ratio will not significantly enhance CO₂ conversion. Moreover, operation in these conditions is questionable, since the separation and recycle of H₂ should be considered. We envision this solution for systems where CO is the main product of interest.

3.4. eRWGS: Effect of GHSV

To further explore the eRWGS at high GHSV, we prepared another washcoated SiSiC foam ($L_{foam} = 8 \text{ cm}$) with a low catalyst density of 13 g/L. This enabled us to operate the system up to $GHSV = 600 \text{ kNl/kg}_{cat}$ h. The washcoated foam had an average catalyst layer thickness of 13.2 μ m, considering that 4/5 of the foam volume was washcoated (Fig. S2). As shown in Fig. 9(a), under kinetic controlled conditions CO_2 conversion decreases on increasing GHSV. Expectedly, such an effect vanishes once equilibrium is reached at the reactor outlet, which occurs at progressively higher temperature on increasing the GHSV. At 700 °C equilibrium conversions were attained, even with the highest space velocity of 600 kNl/kgcat/h. Fig. 9(b) compares the corresponding CH4 outlet fraction results to equilibrium. Hardly any CH4 was detected above 650 °C, consinstently with the equilibrium predictions. At low temperatures, lower CH₄ outlet fractions were obtained with increasing space velocity. Heyl et al. [50], reported that CH₄ was produced with CO as intermediate during CO2 hydrogenation over Rh/Al2O3 catalyst. This indicates that with incrasing space velocity, the methanation reaction becomes slower compared to RWGS, in line with literature reports [31,51].

Fig. 9(c) illustrates the effect of H_2/CO_2 feed ratio on CO_2 conversion at GHSV of 600 kNl/kg_{cat}/h. The system was mainly thermodynamically controlled above 700 °C for all the investigated H_2/CO_2 feed ratios. Fig. 9(d) shows that high H_2/CO_2 feed ratios favored the CH₄ formation at low temperatures, in line with equilibrium.

To verify the stability of the system, eRWGS tests were carried out repeatedly with T_{cap} of 700 °C and 650 °C (H₂/CO₂ = 2, 600 kNl/kg_{cat}/



Fig. 8. CO₂ and H₂ conversions during eRWGS runs with different H₂/CO₂ feed ratios. 1 %Rh/Al₂O₃ washcoated SiSiC foam (L_{foam} = 9.9 cm) with catalyst density of 39.3 g/L. Experiments carried out at 800 °C, GHSV = 100 kNl/kg_{cat}/ h, ambient pressure.



Fig. 9. (a) CO₂ conversion and (b) CH₄ outlet fraction during eRWGS runs with different space velocity: GHSV = 200, 400 and 600 kNl/kg_{cat}/h with a fixed H₂/CO₂ feed ratio of 2; (c) CO₂ conversion and (d) CH₄ outlet fraction from eRWGS runs with different H₂/CO₂ feed ratio of 0.5, 1, 2 and 3 at a fixed GHSV = 600 kNl/kg_{cat}/h. Experiments were carried out with 1 %Rh/Al₂O₃ washcoated SiSiC foam (L_{foam} = 8 cm) with catalyst density of 13 g/L, ambient pressure.

h, ambient pressure) at different working hours (15 h, 60 h, and 75 h), as shown in Fig. 10(a). The two conditions were selected as they are representative of the thermodynamic and kinetic controlled regimes, respectively. In between those selected stability runs, we also carried out eRWGS tests at different conditions, i.e., different space velocities and H_2/CO_2 feed ratios (*c.f.* experimental log from Table S2). We found that the structured catalyst remained stable throughout the testing period, with no significant drop in either CO_2 or H_2 conversions, as shown in Fig. 10(a). Additionally, the electric resistance of the system remained almost constant at 0.32 ohm, indicating the good electric stability of the foam-based reactor. Fig. 10(b) displays photos of the fresh (left) and used (right) SiSiC foam after at least 80 h on eRWGS stream with varied conditions. We observed no significant deterioration of the catalyst layer and just a slight color change (darkening) that can be attributed to Rh reduction.

4. Conclusions

We have proposed and successfully demonstrated the electrified CO₂ reforming of methane (eCRM) and the electrified reverse water-gas shift (eRWGS) reactions for CO2 valorization, based on direct Joule heating of catalytically activated open-cell foams. Such a reactor concept, relying on porous SiSiC foams that act as catalyst supports as well as heating substrates, allows to reach CO₂ conversion approaching equilibrium in a wide range of operating conditions. The eRWGS process exhibits great potential to achieve low specific energy consumptions for CO₂ conversion: calculations showed that 0.7 kWh/Nm³_{CO2} could be reached assuming an overall adiabaticity of 95% and 90% recovery of the sensible heat. Moreover, hardly any CH₄ could be detected in the products during the eRWGS runs above 675 °C. With a H2-rich feed, a CO2 conversion of 94.2% was achieved. Furthermore, the Joule-heated Rh/ Al2O3-coated foam exhibited excellent catalytic and electrical stability for more than 75 h. By replacing fuel combustion heating with Joule heating based on renewable electric energy, both processes are encouraging solutions for CO2 valorization while drastically reducing the carbon footprint for downstream chemical productions. By considering the potential for compact small-scale reactor design as a result of enhanced heat and mass transfer properties, combined with the fast heat transfer dynamics of Joule heating, the proposed electrified CO₂ valorization concept can be regarded as a promising option to mitigate the



Fig. 10. Stability of the washcoated SiSiC foam in eRWGS experiments: (a) CO_2 and H_2 conversions from eRWGS working hours of 15 h, 60 h and 75 h, experiments were carried out at T_{cap} of 700 °C and 650 °C, H_2/CO_2 feed ratio of 2, 600 kNl/kgcat/h, ambient pressure, note that eRWGS runs at varied conditions such as different space velocity, different H_2/CO_2 feed ratio were carried out among those times. (b) Fresh SiSiC foam and used SiSiC foam after at least 80 h on eRWGS stream with different experimental feed conditions. 1%Rh/Al₂O₃ washcoated SiSiC foam ($L_{foam} = 8$ cm) with catalyst density of 13 g/L.

intermittent nature of renewable sources by storing the excess energy in chemicals.

The proposed electrified foam-based reactor system is in principle applicable to the Ni-based catalysts commonly reported for both reforming and reverse-water gas shift reactions. However, the Rh-based catalysts are known to offer higher activity per catalyst volume/mass. Combined with the excellent stability of the catalyst, and with the significantly reduced heat and mass transfer limitations, the Jouleheated Rh-catalyst washcoated foam reactor thus provides significant advantages in terms of process intensification, particularly in view of the design of compact small-scale reactors.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Data availability

Data will be made available on request.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi. org/10.1016/j.cej.2023.143154.

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