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## Introduction

The production of biogas, a mixture containing mainly CH<sub>4</sub> and  $CO_2$ , is a priority in Europe for the provision of sustainable energy. The European energy production derived from biogas was about 190 TWh in 2022 and is expected to double by 2030.<sup>1</sup> The dry reforming of biogas is seen as a sustainable pathway to produce decarbonised hydrogen, although it is accompanied by  $CO_2$  emissions.<sup>2</sup> On the other hand, the cracking of methane can produce CO<sub>2</sub>-free hydrogen, since the carbon is stored in solid form.<sup>3-7</sup> The feedstock of methane can be natural gas (NG) or biogas, which would need to be treated through separation processes to remove CO2, notably to obtain biomethane.<sup>8,9</sup> There are two main approaches for methane cracking: plasma pyrolysis<sup>10</sup> and catalytic pyrolysis.<sup>11-13</sup> However, the thermal decomposition of methane is also being investigated,<sup>14</sup> although the temperatures are much higher and therefore less interesting than with a catalyst. Because of thermodynamic and technological constraints, very high

# Direct biogas reforming to turquoise H<sub>2</sub> and carbon material in a catalytic fluidised-bed reactor<sup>†</sup>

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The conversion of methane or natural gas into turquoise  $H_2$  and carbon represents a promising pathway towards decarbonised energy. The pyrolysis of biogas (mainly a mixture of methane and  $CO_2$ ) is considered to be a negative carbon-emission technology, as the carbon source comes from biomass and the carbon is captured in solid form. In addition, the presence of  $CO_2$  in the gas mixture enables the dry reforming of methane into syngas. In this study, we show that direct biogas conversion in a fluidised catalytic reactor can produce a syngas whose composition meets the requirements for methanol synthesis or for liquid hydrocarbon production *via* the Fischer–Tropsch process. Confirmed by thermodynamic analysis, we show that the reaction proceeds at temperatures above 900 °C to produce carbon materials, whereas at lower temperatures the carbon is converted to CO by the Boudouard reaction. At 950 °C, very high methane and  $CO_2$  conversions are achieved (>90% and 99%, respectively), accompanied by high hydrogen yield (>90%) and the sequestration of carbon into a turbostratic structure. We believe that the direct catalytic reforming of biogas in a fluidised bed, when combined in series with an FTS process, could allow the production of liquid fuels with no need for costly gas separation units, in particular those implemented for  $CO_2$  capture.

methane conversion is difficult to achieve in a steady state. As a result, an  $H_2$  separation unit has to be placed downstream to allow unconverted methane to be recycled and methane to be purified.<sup>15</sup> Hence, from a global process point of view, the pyrolytic conversion of biogas into  $H_2$  would require two major separation steps: an upstream  $CH_4/CO_2$  separation and a downstream  $CH_4/H_2$  separation.

Here we investigate the direct catalytic cracking of biogas, which means that both methane cracking (1) and methane dry reforming (2) are involved.

$$CH_{4(g)} \rightarrow C_{(s)} + 2H_{2(g)} \quad \Delta H_{298K} = 74.9 \text{ kJ mol}^{-1}$$
 (1)

$$CH_{4(g)} + CO_{2(g)} \rightarrow 2CO_{(g)} + 2H_{2(g)} \quad \Delta H_{298K} = 247.3 \text{ kJ mol}^{-1}$$
(2)

This approach faces several challenges: (i) the reaction is heat transport limited due to the endothermicity of both the cracking and dry reforming reactions, (ii) catalyst deactivation, and (iii) the Boudouard reaction (3), which can convert the solid carbon into CO.

$$C_{(s)} + CO_{2(g)} \rightarrow 2CO_{(g)} \quad \Delta H_{298K} = 172.4 \text{ kJ mol}^{-1}$$
 (3)

While methane conversion with stoichiometric or excess amounts of  $CO_2$  (dry reforming) is well known,<sup>16,17</sup> direct

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methane conversion in the presence of  $\mathrm{CO}_2$  has rarely been addressed so far.  $^{18}$ 

The objective of this work was to investigate the direct catalytic cracking of biogas in a fluidised-bed reactor<sup>19–24</sup> to achieve high CH<sub>4</sub> conversion and H<sub>2</sub> yield with carbon storage in a solid form.<sup>25,26</sup> We show that at temperatures above 900 °C, the fluidised bed reactor produces carbon materials, whereas at lower temperatures, the carbon is gasified to CO by the Boudouard reaction. Whereas nickel-based catalysts are the most active catalysts for low-temperature pyrolysis and enable the production of higher-value carbon nanotubes (CNTs),<sup>19,27,28</sup> iron-based catalysts are more appropriate for high reaction temperatures.<sup>29,30</sup> Here we describe the development of an iron@carbon catalyst that can be fluidised for several hours of time on stream without deactivation.

## **Experimental**

#### Thermodynamic study

A thermodynamic equilibrium analysis employing Gibbs free energy minimisation was performed using HSC Chemistry software (version 4.1, Outokumpu Research Oy, Helsinki, Finland). Initial conditions were configured with 0.667 mol of CH<sub>4</sub>, 0.333 mol of CO<sub>2</sub> and 1 mol of C (amorphous), under 1 bar and from 0 °C to 1000 °C. Due to the insensitivity of biogas reforming performance to pressure variations, the analysis was performed under ideal gas conditions, deliberately ignoring the effect of pressure on thermodynamic properties.<sup>31</sup>

#### Catalytic set-up

The fluidised catalytic bed consisted of a quartz tube (ID 16 mm) filled with Fe@carbon catalyst (11 g, h = 5 cm). The synthetic biogas was a mixture of CH<sub>4</sub>:CO<sub>2</sub> in a 2:1 ratio (99.995%, Air Liquide, France). Flow rates were managed using mass flow controllers (F-201CV, Bronkhorst, France). The fluidised-bed quartz reactor was heated with an electrical furnace (TSVH 12/40/305, Elite Thermal Systems Limited, Merket Horborough, United Kingdom). Additional information can be found in ESI† as well as the scheme of the experimental setup (Fig. S1, ESI†).

Two type of experiments were carried out: (i) as a function of temperature, and (ii) as a function of time on stream (TOS). For the former, the temperature was varied between 750 and 950 °C, with 50 °C steps (10 °C min<sup>-1</sup>) and isothermal dwells of 30 minutes during which the catalyst performances were assessed. The reaction was carried out under a pressure of 1 bar with a 300 mL min<sup>-1</sup> total inlet flow rate (CH<sub>4</sub>/CO<sub>2</sub>/N<sub>2</sub> 2:1:1). For the latter study, the reactor was heated to 950 °C in the same conditions as previously. Methane conversion (in the absence of CO<sub>2</sub>) was studied under the same flow and temperature conditions for comparison purposes.

Hydrogen mass balances were calculated for each experiment and were all above 98%. The gas mixtures were analysed online using a micro gas chromatograph (MicroGC 990, Agilent Technologies, France) equipped with two capillary columns: an MS5A SS column (10 m  $\times$  0.25 mm  $\times$  30 µm) for the analysis of H<sub>2</sub>, O<sub>2</sub>, N<sub>2</sub>, CH<sub>4</sub> and CO, and a Poraplot Q UM column (10 m  $\times$  0.25 mm  $\times$  8 µm) for the analysis of CO<sub>2</sub> and hydrocarbons. N<sub>2</sub> was used as internal standard for quantitative analysis. Gases from both columns were analysed with a thermal conductivity detector (TCD).

The weight hourly space velocity (WHSV) was determined by dividing the total flow rate (cm<sup>3</sup> h<sup>-1</sup>) by the catalyst mass (g). Equations defining WHSV are provided in Table S1 (ESI†) for clarity. The conversions (4) and (5), hydrogen yield (6) carbon balance (7) and hydrogen balance (8) of individual gas components were calculated using the following equations:

$$CH_{4 \text{ conversion}}\% = \frac{F_{CH_4}^{\text{in}} - F_{CH_4}^{\text{out}}}{F_{CH_4}^{\text{in}}} \times 100$$
(4)

$$CO_{2 \text{ conversion}}\% = \frac{F_{CO_2}^{\text{in}} - F_{CO_2}^{\text{out}}}{F_{CO_2}^{\text{in}}} \times 100$$
(5)

$$H_{2 \text{ yield}}\% = \frac{F_{H_2}^{\text{out}}}{2\left(F_{CH_4}^{\text{in}}\right)} \times 100$$
 (6)

$$C_{\text{balance}}\% = \frac{F_{\text{CH}_4}^{\text{out}} + F_{\text{CO}_2}^{\text{out}} + F_{\text{CO}}^{\text{out}} + F_{\text{CH}_4}^{\text{out}} + xF_{\text{C}_x\text{H}_y}^{\text{out}}}{F_{\text{CH}_4}^{\text{in}} + F_{\text{CO}_2}^{\text{in}}} \times 100 \quad (7)$$

$$H_{\text{balance}} \% = \frac{4F_{\text{CH}_4}^{\text{out}} + 2F_{\text{H}_2}^{\text{out}} + yF_{\text{C}_x\text{H}_y}^{\text{out}}}{4F_{\text{CH}_4}^{\text{in}}} \times 100$$
(8)

 $F_i^{\text{in}}$  and  $F_i^{\text{out}}$  are the mole flow rates (mol s<sup>-1</sup>) for the component i (CH<sub>4</sub>, CO<sub>2</sub>, H<sub>2</sub>) fed and discharged.  $F_{C_xH_y}^{\text{out}}$  represents the C<sub>2+</sub> hydrocarbons (C  $\geq$  2 atoms) outlet flow (actually negligible).

#### Catalyst preparation

The Fe@carbon catalyst was prepared by a thermal process involving iron powder (Shanghai Laiwu Powder Material Co. Ltd, China) and methane (Air Liquide, France). An amount of 30 g of iron powder was placed in a quartz reactor (16 mm ID) that was heated at 950 °C under CH<sub>4</sub>. After 3 hours under flow, the reaction was stopped, and the material was recovered, lightly crushed and sieved to a size less than 200  $\mu$ m, which allows appropriate fluidisation for the catalytic pyrolysis of biogas.

#### Catalyst characterisation

The C, H, N, (O) and S contents were measured using a Thermo Scientific Flash 2000 series CHNS/O Analyzer equipped with a thermal conductivity detector (TCD). Thermogravimetric analysis (TGA) was carried out in air (30 mL min<sup>-1</sup>) on a Mettler Toledo TGA/DSC 1 STAR<sup>e</sup> System with a 5 °C min<sup>-1</sup> heating ramp. X-ray diffraction analysis of catalysts and recovered carbon materials was carried out on powders using a Bruker D8 Advance A25 diffractometer in Bragg–Brentano configuration equipped with a Cu anode (Cu K $\alpha$  radiation: 0.154184 nm) and a LynxEye 1-D detector. Specific surface areas were measured by nitrogen adsorption at 77 K using a Belsorp Mini X instrument from BEL MicrotracBel Corp. Prior to measurements, materials were degassed at 250  $^{\circ}$ C for one night. The specific surface areas were calculated according to the Brunauer, Emmett and Teller method (BET).

 $\rm H_2O$  vapour adsorption was determined using a Belsorp-Max2 instrument (MicrotracBel) at 293 K. The adsorption temperature was carefully controlled with a thermostatic bath (DyneoDD-300F) with an uncertainty of  $\pm 0.1$  K. Prior to measurements, materials were degassed at 250  $^\circ\rm C$  for one night.

Raman spectra were recorded at ambient temperatures on a LabRAM HR Evolution spectrometer (HORIBA) equipped with a CCD detector. A laser with a wavelength of 532 nm was used for excitation, and the spectra were calibrated by means of the Raman peak of Si at 521 cm<sup>-1</sup>. The acquisition time was adjusted according to the intensity of the Raman scattering.

SEM analyses were carried out with a Zeiss Sigma Field Emission SEM with an Oxford INCA PentaFETx3 EDS system (model 8100). Analyses were conducted on a fresh, uncrushed Fe@carbon catalyst.

## Results and discussion

#### Catalytic results

From 800 to 950 °C, the fluidised bed exhibited pulsating behaviour within the column, suggesting a bubbling regime (see movie in ESI<sup>†</sup>). This effect is attributed to the temperature increase, leading to the expansion of introduced gases, greater production of reactive gases and, consequently, bubble formation in the fluidised bed. This behaviour is intensified by the use of small catalyst particles with a pseudo-uniform distribution in the reactor and also by the production of carbon. The accumulation of carbon could promote different hydrodynamic resistances, influencing the fluidisation pattern and contributing to the piston effect. Moreover, the fluidisation pattern is markedly influenced by the gas flow rate and the minimum fluidisation velocity of the system.<sup>19,32</sup> It is observed that when the gas flow rate exceeds the minimum fluidisation velocity, bubbles can form and grow with increasing ratio.<sup>33</sup> The observed fluidisation looks very similar to that which is visible in the video made by Dadsetan et al.,26 who, however, used activated carbon as catalysts.

At 750 °C, methane conversion is negligible and increases rapidly as the temperature increases. The CO<sub>2</sub> conversion increases with temperature also, but remains high (>80%) regardless of temperature due to sub-stoichiometric CO<sub>2</sub> in the inlet mixture. At 950 °C, the conversion of methane and CO<sub>2</sub> achieves maximum levels of 91 and 99%, respectively (Fig. 1).

We can also observe that after 150 min, when the temperature is decreased, the conversion level is higher than that obtained at the same temperature at the beginning of the experiment, which indicates an activation phenomenon with time on stream. This is particularly visible at 800  $^{\circ}$ C, for which



Fig. 1 (a) Methane and CO<sub>2</sub> conversion as a function of temperature, (b) carbon balance estimated from gas-phase reactants and products. Carbon balance values above 100% indicate carbon loss from the catalyst; values below 100% indicate an accumulation of solid carbon. Inlet flow: 300 mL min<sup>-1</sup> CH<sub>4</sub>/CO<sub>2</sub>/N<sub>2</sub> 2:1:1.

the conversion is 20% when the temperature ramps up and 30% when it ramps down.

The carbon balance is calculated from a quantitative analysis of the gaseous reactants and products. Hence, a carbon balance below 100% implies that a lesser quantity of C-containing compounds is found at the reactor outlet in the gas phase than in the inlet feed, *i.e.*, solid carbon is formed. Conversely, when the carbon balance is above 100%, a greater quantity of C-containing gas-phase compounds is found at the reactor outlet than in the inlet feed, which corresponds here to the gasification of solid carbon into CO. The evolution of the carbon balance as a function of temperature clearly indicates carbon loss, *i.e.*, production of  $CO_{(g)}$  from solid carbon below 850 °C and carbon formation from gaseous reactants above 850 °C.

Methane conversions in biogas as a function of temperature are slightly higher than the conversion of pure methane at similar temperatures (Fig. 2). Actually, dry methane reforming (CH<sub>4</sub> + CO<sub>2</sub>) occurs during the reaction, resulting in higher methane conversion in the presence of CO<sub>2</sub>. In pure methane cracking (Fig. S2, ESI<sup>†</sup>), the carbon balance never exceeds 100%, indicating that carbon is not gasified in the absence of CO<sub>2</sub>. As the temperature increases, the carbon balance

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**Fig. 2** Comparison of carbon balance (%) and CH<sub>4</sub> conversion (%) for biogas reforming (CH<sub>4</sub>/CO<sub>2</sub>/N<sub>2</sub> 2:1:1) and methane cracking (CH<sub>4</sub>/N<sub>2</sub> 1:1) at T = 950 °C, 1 bar, 300 mL min<sup>-1</sup>. A carbon balance of less than 100% (grey dotted line) indicates carbon storage, while above 100% there is carbon gasification.

decreases to a minimum of 17% at 950  $^\circ C$ , indicating that most of the methane is converted to solid carbon on the catalyst.

Hydrogen yield (Fig. 3a) as a function of temperature follows the profile of methane conversion, reaching a maximum yield of 91% at 950 °C. The H<sub>2</sub> productivity increases with the temperature and reaches about 950 mmol<sub>H2</sub>  $g_{cat}^{-1}$  min<sup>-1</sup> for this yield. C2 and C3 hydrocarbons are observed in trace concentrations.

The evolution of the  $H_2/CO$  ratio as a function of  $H_2$  yield is shown in Fig. 3b. Interestingly, as the temperature increases,



Fig. 3 (a) Hydrogen yield (%) and productivity  $(mmol_{H_2} g_{cat}^{-1} min^{-1})$  over time and (b) H<sub>2</sub>/CO ratio as a function of H<sub>2</sub> yield (%).

the  $H_2/CO$  ratio increases linearly with the  $H_2$  yield. The same trend is observed in the thermodynamic study (Fig. 4). Compared to the  $H_2$  productivity, the CO productivity is less affected by temperature. This can be explained by the ability of the carbon from the catalyst to shift the equilibrium of the Boudouard reaction, increasing the CO concentration in the reaction and reversing the trend.

The Fe@carbon catalyst also showed stability over for 2 hours on stream at 950 °C with biogas (Fig. S3, ESI<sup>+</sup>) and more than 3h for pure methane (Fig. S4, ESI<sup>+</sup>). The methane conversion showed a slight activation in the first tens of minutes, then was stable and progressively decreases after 2 h on stream, accompanied by an increase in carbon balance (Fig. S3a, ESI<sup>†</sup>), as observed by Lim *et al.*<sup>34</sup> At long TOS, less H<sub>2</sub> was produced, while CO production remained constant. During this long time-on-stream test, a mass gain of 9 g was observed (carbon storage) and the catalytic bed volume was four times higher than for the fresh catalyst (Fig. S5, ESI<sup>†</sup>). The apparent density of the Fe@carbon catalyst decreased from 1.22 g cm<sup>-3</sup> (fresh) to 0.38 g cm<sup>-3</sup> (spent). As the reaction proceeds, some lighter catalyst particles (1.94 g) escape from the top of the fluidised bed which could explain in part the decrease of methane conversion. Several studies have shown that the catalyst can be regenerated with different gas streams such as CO<sub>2</sub>, air and H<sub>2</sub>O vapor.<sup>32</sup> This could prevent catalyst deactivation over a TOS reaction period. Under the same WHSV and temperature conditions, the catalyst developed by Lim et al.34 shows lower CO<sub>2</sub> and CH<sub>4</sub> conversions. In other studies, notably the work of Torres et al.,33 an iron-based catalyst has been tested under similar conditions to ours (e.g. WHSV and temperature) and presents a lower activity range.

The fluidisation method possesses significant advantages by circumventing the inherent heat transport limitations often encountered in traditional reaction systems. A detailed study of fluidised systems in comparison to others is given elsewhere.<sup>35</sup> These include the superior temperature uniformity throughout the reactor, thereby eliminating the occurrence of hot spots and avoid the reactor plugging during the carbon production. However, operating at temperatures as high as 950 °C presents challenges as catalysts can sinter and prevent effective fluidisation. Tackling these constraints is critical to optimizing the



Fig. 4 Thermodynamic equilibrium for biogas decomposition as a function of temperature. Conditions:  $CH_4 = 0.667$  mol,  $CO_2 = 0.333$  mol, C(A) = 1.000 mol, 1 bar.

performance and efficiency of this process in future research and scale-up efforts.

#### Thermodynamic aspects

Methane cracking and biogas reforming are endothermic equilibrated reactions, so increasing the reaction temperature shifts the equilibrium towards the production of H<sub>2</sub>. In our case, we should keep in mind that the fresh Fe@carbon catalyst contains about 20% of its weight in carbon (Fig. S6, ESI<sup>†</sup>). Hence, under certain conditions it can produce CO by carbon or coke gasification (with  $CO_2$ ), while under other conditions it can accumulate solid carbon deposits, most likely by coking of the iron surface. Below 850 °C, the carbon previously deposited onto the iron is gasified to CO by the Boudouard reaction  $(C_{(s)} +$  $CO_{2(g)} \rightarrow 2CO_{(g)}$  (Fig. 1). This indicates that carbon formation by methane cracking  $(CH_{4(g)} \rightarrow C_{(s)} + 2H_{2(g)})$  is slower than carbon gasification by the Boudouard reaction. In contrast, at temperatures above 850 °C, solid carbon is produced, because the methane cracking reaction produces more carbon than does gasification by the Boudouard reaction. These results are consistent with the thermodynamic study (Fig. 4). Indeed, the thermodynamic equilibrium is achieved by considering the carbon on the catalyst, denoted C(A) for amorphous carbon.

In this equilibrium system, the carbon composition decreases from 500 °C onwards, in favour of CO (gasification of carbon with CO<sub>2</sub> *via* the Boudouard reaction). Beyond 650 °C, the amount of carbon increases, while the amount of CH<sub>4</sub> decreases. Above 900 °C, thermodynamic calculations show that the concentrations of CH<sub>4</sub>, CO<sub>2</sub> and water vanish and the composition reaches a plateau consisting of 1.15 mol of H<sub>2</sub>, 0.66 mol of CO and 1.24 mol of solid carbon. The experimental conversions at 900–950 °C match with thermodynamic estimates, especially the carbon yield (Fig. 1). Also, the experimental H<sub>2</sub>/CO ratio is 1.9:1 at 950 °C, while the thermodynamic estimate is 1.7:1 (Fig. 4). Thus, we can confirm that at 950 °C the thermodynamic equilibrium is reached with a H<sub>2</sub>: CO ratio close to 2, which is ideal for FTS, MeOH and DME synthesis,<sup>36,37</sup> while the carbon is stored in solid form.

Our results at high temperature are in agreement with the work of Li *et al.*<sup>18</sup> As a matter of fact, the CH<sub>4</sub> and CO<sub>2</sub> conversions obtained are very close after 900 °C. That said, the H<sub>2</sub>/CO ratio is higher in our case.

#### Catalyst and carbon product characterisation

TGA of the fresh Fe@carbon, carried out in air between 25 and 1000  $^{\circ}$ C (5  $^{\circ}$ C min<sup>-1</sup>), shows that the catalyst contains 20 wt% carbon and 74 wt% of a mixture of iron and iron oxide (Fig. S6, ESI†). A slight increase in mass can be seen at 500  $^{\circ}$ C (followed by another around 750–900  $^{\circ}$ C), corresponding to the different stages of Fe oxidation.<sup>38</sup> The carbon on the catalyst has a decomposition temperature under air of around 630  $^{\circ}$ C, corresponding to a form of pyrolytic graphitic carbon. By comparing the TGA of the fresh and spent catalysts, we can see that the amount of carbon has increased from a mass of 20% to a mass of 74%.

XRD analysis of fresh and spent catalysts shows several characteristic graphite lines (Fig. S7, ESI†), including the main



Fig. 5 SEM picture of the (a) fresh Fe/C catalyst and (b) spent Fe/C catalyst.

graphite line at 26.6°. The characteristic lines of metallic iron can also be seen, with the most intense line at  $44.7^{\circ}$ . By comparing the diffractograms of the fresh and spent catalysts, we can see that the intensity of the characteristic graphite line on the catalyst increases after the test.

Raman spectroscopy is a relevant technique for characterising local orders in carbonaceous materials, thanks to the  $I_{\rm D}/I_{\rm G}$ ratio (Fig. S8, ESI<sup> $\dagger$ </sup>). The G band (1580 cm<sup>-1</sup>) represents the vibrational mode associated with the asymmetric stretching of all sp<sup>2</sup> carbon atoms, while the D band is attributed to the breathing mode in the aromatic rings of Poly Aromatic Hydrocarbons (PAHs).<sup>39,40</sup> The presence of the D band ( $1350 \text{ cm}^{-1}$ ) in these materials is attributed to structural defects. It has been shown that the intensity of the D band relative to that of the G band increases with the amount of disorder.<sup>41-43</sup> In this study, the carbon deposits before and after biogas cracking present the same  $I_{\rm D}/I_{\rm G}$  ratio ( $\approx 0.42$ ). This ratio is similar to the one determined by Charisiou et al.44 This indicates that the two different carbon deposits have the same local order or degree of crystallinity which correspond to turbostratic carbon class. The spent catalyst has a specific surface area (SSA) of 11 m<sup>2</sup> g<sup>-1</sup>, compared to 2 m<sup>2</sup> g<sup>-1</sup> for the fresh catalyst (with a pore volume

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of 26.6 cm<sup>3</sup> g<sup>-1</sup> and 6.8 cm<sup>3</sup> g<sup>-1</sup>, respectively)(Table S2, ESI†). This additional surface and pore volume is caused by carbon deposition during biogas conversion. (Fig. S9, ESI†). H<sub>2</sub>O physisorption on the fresh and spent catalyst clearly indicates that the two samples are hydrophobic (Fig. S10, ESI†) and adsorb very low amount of water (Table S3, ESI†). However, Fe/C spent catalyst presented a higher amount than fresh Fe/C catalyst, and can be explain because of the higher surface area of the catalyst determined by N<sub>2</sub> physisorption.

SEM was used to study the morphology of the fresh and spend Fe@carbon (Fig. 5). The carbon particles on the fresh catalyst appear as in the form of thin, smooth plates, similar to the findings of Rew *et al.*<sup>45</sup> The particles of spent catalyst after biogas reforming appear as carbon filaments with heavier particles on their ends corresponding to iron particles, which may give an indication of Carbon Nanotubes (CNTs) formation during the biogas reforming. However, the quantity of CNT is too small to be detected by Raman spectroscopy between 120 and 350 cm<sup>-1</sup> as shown by the work of Dresselhaus *et al.*<sup>46</sup>

## Conclusions

In this work we have demonstrated the development of a Fe@carbon catalyst that is very easy to manufacture and that can be used as a fluidised catalyst for biogas pyrolysis. Catalytic performances are very stable for at least 2 hours of TOS. We show that in the catalytic conversion of biogas, temperatures in excess of 850 °C are required for carbon production; otherwise, the carbon is gasified into CO. Compared to pure methane cracking, biogas reforming can be achieved with very high methane conversion (90% vs. 83%) at 950 °C, thanks to the additional dry reforming reaction. Equally important is the H<sub>2</sub>:CO ratio approaching 2:1, an ideal ratio for Fisher-Tropsch or MeOH synthesis. As a result, this technology would allow the production of a syngas directly from biogas with an integrated carbon capture in the form of solid carbon. Indeed, at 950 °C, the outlet gas concentration would be  $H_2$  (62%) and CO (33%), with lower concentrations in  $CH_4$  (4%) and  $CO_2$  (1%), while the carbon production would be 0.3 kg<sup>-1</sup> kg<sub>catalyst</sub><sup>-1</sup> h<sup>-1</sup>.

This work opens up the prospect of converting biogas into liquid hydrocarbons in two catalytic steps, without the need for gas separation (*e.g.*,  $CH_4/CO_2$ ) or recycling ( $CH_4/H_2$ ) at low expenditure and environmental costs, taking also advantage of the availability and non-toxicity of iron ores. As part of our EU grant program (TITAN), we plan to scale up the reactor for a comprehensive techno-economic analysis, assessing  $H_2$  production costs, greenhouse gas emissions and environmental impacts to guide future industrial implementation and sustainable energy innovation.

## Author contributions

Valentin L'hospital: investigation, validation, visualization, writing – original draft. Leandro Goulart de Araujo: writing – review & editing. Yves Schuurman: writing – review & editing. Nolven Guilhaume: project administration, writing – review & editing. David Farrusseng: supervision, project administration, writing – review & editing,

## Conflicts of interest

There are no conflicts to declare.

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