Thermodynamic analysis of an improved integrated biomass gasifier solid oxide fuel cell micro combined heat and power system

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A B S T R A C T

Limited overall efficiency and excessive complexity can hinder the competitiveness of biomass gasifier solid oxide fuel cell micro combined heat and power systems. To overcome these problems, hydrocarbons direct internal reforming is analysed as a strategy to increase efficiency and reduce system complexity. To the same end, two biosyngas heating-up strategies are compared: catalytic partial oxidation and afterburner off-gases utilization. A comprehensive approach combining thermodynamic equilibrium calculations, experimental measurements, and system modelling was used. The gas cleaning unit should operate at 400 °C to decrease H₂S and HCl below 1 ppmv. A tar amount of 120–130 g Nm⁻³ dry biosyngas for woodchips and 190 g Nm⁻³ for straw pellets was measured and 2-methoxyphenol, hydroxyacetic acid and hydroxyacetone were selected as representative compounds. With direct internal reforming the cathode air flow rate decreases from approximately 90 kg h⁻¹ to 60 kg h⁻¹. This leads to an increase of around 1% point in electrical efficiency and of even 5–6% points in thermal efficiency. Direct internal tar reforming seems therefore an advantageous strategy. The catalytic partial oxidation unit increases the system overall efficiency but reduces the electric efficiency from roughly 38%–30% and is therefore not advised.

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1. Introduction

Combining biomass gasification with high temperature fuel cells has received considerable attention as an alternative to fossil fuel based energy systems [1–4]. In a biomass gasifier solid oxide fuel cell (SOFC) system, the biomass is converted into biosyngas, a mixture of hydrogen, carbon monoxide, carbon dioxide, nitrogen, and methane, which is converted into heat and electricity in the SOFC. Biosyngas also contains minor compounds which are harmful for downstream equipment and have to be removed: sulphur, halides, particulate matter and tar compounds [5]. The biosyngas composition and the content of tar are highly dependent on the gasification agent used, that can be air, oxygen, or steam [6]. Before entering the SOFC, the gas passes through a gas cleaning unit (GCU) where the contaminants are removed at low (<300 °C) or high temperature. Gas cleaning is a crucial step in the system heat management, and it strongly affects the complexity and overall system efficiency.

In literature, system modelling has been used to determine the obtainable electric and overall efficiencies with the different cleaning methods, and to optimize the design of biomass gasifier SOFC systems. Omosun et al. modelled two combined heat and power (CHP) systems: a downdraft gasifier with cold gas cleaning, obtaining an efficiency of 20.8% electrical and 33.9% overall, and a fluidized bed gasifier with hot gas cleaning, achieving an efficiency of 22.6% electrical and 59.6% overall [7]. Aravind et al. evaluated the performance of a 100 kW gasifier SOFC system with a gas turbine (GT) bottoming cycle. With gas cleaning at 750 °C, the system achieved 54% electrical and 77–78% total efficiency, whereas cleaning the gas at ambient temperature resulted in 49.5% electrical and 49.9–57% total efficiency [8]. Toonssen et al. investigated the performance of a 30 MWₑ integrated gasifier hybrid SOFC-GT systems. Cleaning the gas at 700 °C instead of 120 °C gave an increase in the efficiency of roughly 1% [9]. Liu et al. found that cleaning the gas at 450 °C gave a CHP energy efficiency of 65%, while cleaning the gas at ambient temperature lowered the efficiency to 57% [10]. More recently, Doherty et al. found that in a 120 kWₑ, dual fluidized bed steam gasifier SOFC system, increasing the cleaning temperature from 400 °C to 700 °C increased the system net electrical efficiency from 25.3% to 30.2% but decreased the CHP
efficiency from 69.5% to 67.0% [11]. Several recent works in the literature discuss other important aspects of biomass gasifier solid oxide fuel cell systems, or their integration with additional bottoming cycles [12–15].

Among biosyngas contaminants, tar, together with light condensable and volatile organic compounds, require particular attention. These compounds can condense in cold spots and cause carbon deposition in downstream equipment. However, they are hydrocarbons and could be regarded as useful fuel. In low-temperature GCUs, tar, light condensable and part of the volatile organic compounds are removed from the gas. Differently, in high-temperature GCUs, tar, light condensable and part of the volatile organic compounds could be regarded as useful fuel. In low-temperature GCUs, tar, light condensable and part of the volatile organic compounds are removed from the gas. Differently, in high-temperature GCUs, these compounds are converted into H2 and CO by reforming in a dedicated reactor via the endothermic reactions (1–3) [16,17].

Steam reforming: $\text{C}_x\text{H}_y\text{O}_z + (x - z)\text{H}_2\text{O} \rightarrow x\text{CO} + [(x + y/2 - z)]\text{H}_2$

(1)

$\text{C}_x\text{H}_y\text{O}_z + n\text{H}_2\text{O} \rightarrow (n + x/2)\text{H}_2 + n\text{CO}$

(2)

Dry reforming: $\text{C}_x\text{H}_y\text{O}_z + n\text{CO}_2 \rightarrow (x/2)\text{H}_2 + 2n\text{CO}$

(3)

The heat required for these reactions can be provided by the SOFC outlet gas, either via heat exchangers or anode recirculation, or by biosyngas partial oxidation. However, heat transfer limitations decrease the overall system efficiency, while partial oxidation and recirculation lower biosyngas reactant concentrations. Reforming might take place internally in the SOFC due to the presence of Ni catalyst and the high operating temperature. In internal reforming, the heat required is provided directly by the SOFC, and the reactions can take place in a separate chamber (indirect internal reforming) or in the anode chamber (direct internal reforming), as illustrated in Fig. 1. Internal reforming eliminates the need of an additional reactor in the GCU, and improves and simplifies the system heat management. Moreover, the endothermic reforming reactions can cool down the SOFC, thus decreasing the energy consumed by the cathode air blower and making the system more efficient.

Some studies have already analysed the advantages of internal CH4 reforming in high temperature fuel cell systems. Lanzini et al. investigated the efficiency and economic gain obtainable by increasing syngas CH4 content in an integrated coal gasifier SOFC system [18]. In a previous study, Nagel et al. compared seven 1 MWc integrated biomass gasifier SOFC systems. Two configurations adopted internal CH4 reforming and a methanation reactor to increase CH4 amount in biosyngas [19,20]. Di Carlo et al. compared the efficiency of a molten carbonate fuel cell coupled with a fast internally circulated fluidized-bed biomass gasifier with internal or external methane reforming; the former increased the stack efficiency from 42% to 53% and reduced the cathode flow rate by 30% [21]. In the analysis of a 120 kWc biomass gasifier SOFC and Stirling engine system, Rokni concluded that using a methanizer before the SOFC reduced the cathode air necessary for cooling, hence less SOFC outlet gas heat was necessary to preheat biosyngas, thus increasing the system electrical efficiency by 2% points [22]. While the benefits of CH4 internal reforming have been investigated, little if any information is available on the advantages of direct internal reforming of other hydrocarbons (i.e., tar, light condensable and volatile organic compounds).

The competitiveness of biomass gasifier SOFC systems, especially at micro scale (<250 kW fuel input power), can be hindered by the limited overall system efficiency typical of low temperature gas cleaning units and by the complexity of both low and high temperature GCUs. To overcome these problems, within the “FlexiFuel-SOFC” project, TU Delft investigated the possibility to adopt new design strategies. In this work, we use a comprehensive approach combining thermodynamic equilibrium calculations, experimental measurements, and system modelling to evaluate the efficiency gain obtainable with direct internal reforming of hydrocarbons. Moreover, we compare two strategies to heat up the biosyngas leaving the GCU and entering the SOFC: a catalytic partial oxidation reactor where part of the biosyngas is combusted, and a heat exchanger where the SOFC afterburner off gases are used. The analysis covers two biomass feedstocks with different humidity content since this parameter affects significantly the biosyngas composition and the tar content. An updraft gasifier was selected based on a. the possibility to handle different types of biomass in terms of moisture and size, b. the low amounts of particulate matter in the biosyngas, c. the high gasification efficiency [23]. Furthermore, the high quantity of light condensable and volatile organic compounds and tar produced in an updraft gasifier (around 150 g Nm$^{-3}$ dry basis [23]) presents an opportunity for direct internal tar reforming. The results of the study are expected to contribute to the development of micro scale biomass gasifier fuel cell systems.

2. System configuration

The system in analysis is being developed under the European
3. Methodology

Afterburner off gases. All the cases analysed, the cathode inlet air is pre-heated by the AVL. The system includes an afterburner where the residual fuel in different compositions is extracted from the gasifier in a slip stream and, after the contaminant removal stage, is fed to the SOFC. The focus of the study is on the SOFC system and on the gas cleaning step; therefore, the model does not include the gasifier, but it starts from the slip stream and ends with the boiler (part of the gasifier), where the SOFC afterburner off gases are used to generate heat together with the remaining biosyngas not extracted for electricity production. The gasifier (PuroWIN technology), including the burner and the boiler, was developed by WindHager in cooperation with the research institute BIOS Bioenergysysteme.

The GCU, developed by TU Delft and HyGEAR, has to provide clean the biosyngas from sulphur and chlorine and a ceramic filter for particulate matter removal. For what concerns tar, volatile organic and light condensable compounds reforming, and the heating up of the biosyngas between the GCU and the SOFC, four different configurations were analysed:

1. CASE A: External reforming - Heat in the afterburner off gases is used for the reforming reactions and to heat up the biosyngas entering the reformer.
2. CASE B: External reforming - Heat in the afterburner off gases is used for the reforming reactions. A catalytic partial oxidation (CPOX) reactor is used to heat up the biosyngas entering the reformer.
3. CASE C: Internal reforming – A CPOX reactor is used to heat up the biosyngas entering the SOFC.
4. CASE D: Internal reforming - Heat in the afterburner off gases is used to heat up the biosyngas entering the SOFC.

The SOFC stack system was developed by Fraunhofer IKTS and AVL. The system includes an afterburner where the residual fuel in the anode outlet gases is combusted with the cathode outlet air. In all the cases analysed, the cathode inlet air is pre-heated by the afterburner off gases.

3. Methodology

To evaluate the system efficiency gain obtainable with direct internal reforming and identify the most efficient biosyngas heating-up strategy, a comprehensive approach combining system modelling, thermodynamic equilibrium calculations and experimental data from the gasifier of the “FlexiFuel-SOFC” project was used. In this section, each activity is described in detail.

3.1. Thermodynamic equilibrium calculations

Thermodynamic equilibrium calculations were used first in the system conceptual design phase to compute the steam necessary to avoid carbon formation, and second to discover which sulphur and chloride compounds can be expected in the biosyngas, thus helping with the selection of suitable sorbents. Finally, the calculations were used to determine the GCU operating temperature allowing to achieve the target cleaning values. The software FactSage v5.4.1 was used for the thermodynamic equilibrium calculations. Taking as inputs the mass of the reactants, process temperature, and pressure, the software gives as outputs the products and their amount based on Gibbs Free Energy minimization [25]. To calculate the necessary steam flow rate, the input mass of reactants used was the gas composition arriving from the gasifier and the temperature was set to 350 °C, that is the lowest operating temperature considered for the GCU.

The main gas composition from the gasifier was taken from computational fluid dynamics (CFD) simulations performed by BIOS applying in-house developed and experimentally validated models. Nearly steam-saturated air at 75 °C was used as gasifying agent, and two feedstocks, woodchips (moisture content 30% in weight) and straw pellets (moisture content 8% in weight), were studied [26–29]. While the main gas composition was obtained from simulations, the amounts of tar, light condensable and volatile organic compounds were measured during test runs. More details on the measurement procedure are given in section 3.2.

The amounts of sulphur and chlorine in the biosyngas extracted with the slip stream were obtained from a mass balance. The inlet amount was measured with an ultimate analysis of the feedstock. The outlet amount was divided in three parts: one part in the gasifier ashes, one part in the flue gases from the gasifier, and one part in the slip stream biosyngas. The amounts in the ashes and in the flue gases were measured, while the amount in the slip stream biosyngas was obtained as difference. A sulphur content of 35.3 and 357.1 mg kg⁻¹ humidified biosyngas was calculated for woodchips and straw pellets, respectively. Regarding chlorine, a concentration of 13.3 and 1618.2 mg kg⁻¹ humidified biosyngas was calculated. High temperature sorbents able to remove the S and Cl compounds
present at equilibrium at 400 °C were selected from literature. The equilibrium concentrations of gaseous sulphur and chlorine compounds at the outlet of the GCU were calculated in the temperature interval 350–450 °C using 1 kg of biosyngas and 10 kg of the identified sorbents. The maximum temperature meeting the SOFC requirements was chosen as the GCU operating temperature.

3.2. Experimental data: tar sampling and analysis

To develop a system model as close as possible to the real system, an experimental campaign was conducted to sample and analyse the quantity and the tar species generated in the updraft gasifier. This approach was deemed very important for the study since the heat required to reform tar is highly dependent on the amount and chemical composition of tar and this is strictly dependent on the biomass used, the type of gasifier and the operating conditions [30].

The gasifier was operated two days with woodchips and one day with straw pellets. While wet sampling was used to measure tar compounds, the amount of some light hydrocarbons (propene) and some light condensable species (formaldehyde, acetic acid, methanol and ethanol) were measured with Fourier-transform infrared spectroscopy (FT-IR). The sampling was performed following the tar protocol (CEN TC BT/TF 143 WI CSC 03002.4) [31]. The samples were analysed with gravimetric analysis in BIO5 Bioenergysysteme, Graz, and with gas chromatography mass spectrometry (GC-MS) analysis in Energy Research Center (ECN) of the Netherlands, Petten. In gravimetric analysis, the samples were evaporated for roughly 20 h at ambient pressure and 60 °C. An elemental analysis was done on the gravimetric tar to determine the weight fractions of carbon, oxygen, hydrogen and nitrogen. In GC-MS analysis, the GC oven was operated between 40 °C and 245 °C and GC method B.C302 (suitable for pyrolysis oil type tar) was used. Gravimetric analysis gives the amount of condensable tar present in syngas but not their chemical composition. Moreover, during this analysis, some of the volatile organic and light condensable compounds are lost thus giving an incomplete picture of the biosyngas composition. Conversely, GC-MS analysis can measure light condensable and volatile organic compounds, and gives the amounts of single tar compounds present in the mixture. Unfortunately, GC-MS cannot detect very heavy tar (Class 1) which are complex poly aromatic hydrocarbons, or complex molecular structures similar to the one found in pyrolysis oil and often referred to as pyrolytic lignin. In this work, it was assumed that all the compounds present in gravimetric tar could be represented by a selection of a few representative primary tar compounds.

An empirical formula for gravimetric tar was calculated from the gravimetric tar elemental analysis. Two of the most abundant compounds from GC-MS analysis were selected and, together with a third compound selected among primary tar compounds reported in Ref. [30], their weight percentage in biosyngas was calculated to match with the empirical formula calculated. These three compounds were used to represent gravimetric tar. The compounds selected have boiling points at atmospheric pressure higher than 60 °C and vapour pressure at 25 °C lower than that of isopropanol. These constraints were chosen to avoid selecting compounds which are not actually present in gravimetric tar.

3.3. System modelling

A system model was created using the software Aspen Plus, as illustrated in Fig. 2.

In all the simulated cases, the fuel utilization and the electricity produced by the SOFC (SOFC-DC) remained constant and equal to 0.8 and 6 kW, and the biosyngas flowrate required (SLIP-GAS) was calculated accordingly. The biosyngas is assumed to be extracted from the gasifier at 440 °C, and is then humidified with superheated steam (SLIP-WT) at 260 °C. In the real system, the heat from the combustion of the remaining biosyngas is used to generate the required steam.

The HCl, H2S and particulate matter removal stages of the GCU are modelled with two heat exchangers. The first accounts for the heat losses to the environment, and the second one for the heat transfer from the afterburner (AFTBRN) off gases. The heat losses are assumed equal to 400 W and were calculated from a preliminary coarse design of the gas cleaning unit based on previous experience in the “BioCellus” project [32]. The biosyngas leaves the GCU at 395 °C. In case A, biosyngas is heated up to 740 °C in the tar reformer (TAR-REF). At this operating temperature, tar components are expected to be reformed [33]. The tar reformer is modelled with a Gibbs reactor with heat losses to the environment equal to 160 W. Also in this case, the heat losses were assumed based on a preliminary coarse design of the component. In cases B and C, biosyngas enters the CPOX unit where it is partially combusted. The unit is modelled with a stoichiometric reactor with heat losses to the environment of 160 W assumed equal to those of the tar reformer. The air flow rate is calculated in order to reach a biosyngas temperature of 740 °C. In case D, biosyngas enters a heat exchanger (ANODE-HX) where it is heated up to a temperature of 740 °C by the afterburner off gases. Table 1 summarizes the four cases analysed.

The software Aspen Plus does not include an SOFC model. Therefore, the SOFC is modelled using four blocks: a Gibbs reactor (ANODE) that simulates the electrochemical reactions; a separator (CA-SEP) that separates the amount of oxygen ions crossing through the electrolyte: a heat exchanger (CA-HEAT) to calculate the amount of air required to maintain the SOFC temperature; and a splitter to split the electricity and heat losses from the total enthralpvy variation in the anode reactor. In this SOFC model, the fuel utilization, cell active area and area specific resistance are given as input parameters. The heat losses of the SOFC (SOFC-HLS) are assumed equal to 900 W. The values were taken from the stack manufacturer and from previous experience of AVL. The equations used to model the SOFC and to perform the electrochemical calculations are presented in Appendix A.1. Air at 25 °C is preheated in a heat exchanger (AIR-HX) by the exhaust gases coming from the afterburner up to 700 °C before entering the SOFC. This value was obtained by simulations performed by AVL using suppliers data to model the heat exchanger.

The outlet flows from the SOFC are fed into the afterburner, modelled with a stoichiometric reactor with heat losses of 220 W. The afterburner off gases are then separated: one part is used to heat the cathode air and the other part is used for biosyngas heat requirements. The two streams are then merged again before being fed to the heat recovery section (HT-RECOV) where hot water at 80 °C is produced. The exhaust flow at 84 °C is finally compressed in a blower (SOFC-CMP) and discarded to the environment at 133 °C (SOFC-FG).

A power electronics unit is used to condition the electricity produced in the SOFC (W-CDTNR). This unit is composed of two components: a DC/DC rectifier and a DC/AC converter. Part of the electricity (W-COMP) is supplied to the SOFC blower, part is lost due to conversion losses (EL-LOSS), and the remaining is converted into useful AC power (W-NET). Table 2 summarizes the input values used in the model. Where not previously specified, values were taken from prior experience of project partners.
4. Results

4.1. Thermodynamic equilibrium calculations

Equilibrium calculation showed that at 350 °C, which is the lowest temperature considered for the GCU operation, the steam amount necessary to avoid carbon formation is 0.12 and 0.25 kg per kg of biosyngas in case of woodchips and straw pellets, respectively. Table 3 presents the gas composition after the steam addition.

The calculations also showed that at 400 °C, S and Cl are mostly present as HCl, H2S, and to a minor extent as COS, as shown in Table 4, where the contaminants concentrations are reported. For both feedstock, the concentration of contaminants in biosyngas is above the SOFC thresholds of 1 ppmv for H2S and 5 ppmv for HCl. The biosyngas derived from straw pellets contains a significantly higher concentration of HCl (two orders of magnitude) and also of H2S (one order of magnitude).

Alkaline earth and alkali metal-based sorbents can be used to decrease the concentration of HCl [35]. The reactions are illustrated as follows:

\[ 2\text{HCl} + 2\text{NaAlO}_2 \rightarrow 2\text{NaCl} + \text{Al}_2\text{O}_3 + \text{H}_2\text{O} \]  \hspace{1cm} (4)

\[ 2\text{HCl} + \text{Na}_2\text{CO}_3 \rightarrow 2\text{NaCl} + \text{CO}_2 + \text{H}_2\text{O} \]  \hspace{1cm} (5)

Potassium compounds are capable of achieving lower residual HCl concentrations as compared to sodium compounds. However, the vapour pressure of KCl is higher than that of NaCl, the two products of the cleaning reactions [36]. Nonetheless, potassium carbonate was selected as sorbent for HCl removal since the operating temperature is too low to result in relevant concentrations of gaseous KCl. The equilibrium HCl concentration is roughly two orders of magnitude lower than the limit value in the whole temperature range (see Table 5). The results are in agreement with the calculations and experiments performed by Krishnan et al. [36,38].

Metal based sorbents, such as zinc or copper oxides are a good option for H2S removal from biosyngas, according to reactions (6) and (7) [35]. Moreover, carbonyl sulfide might be also directly removed by these sorbents, or it can be converted to H2S in the GCU according to the hydrolysis reaction (8) [34].

\[ \text{H}_2\text{S}(g) + \text{ZnO}(s) \rightarrow \text{ZnS}(s) + \text{H}_2\text{O}(g) \]  \hspace{1cm} (6)

\[ \text{H}_2\text{S}(g) + \text{CuO}(s) \rightarrow \text{CuS}(s) + \text{H}_2\text{O}(g) \]  \hspace{1cm} (7)

\[ \text{COS}(g) + \text{H}_2\text{O}(g) \rightarrow \text{H}_2\text{S}(g) + \text{CO}_2 \]  \hspace{1cm} (8)

ZnO was selected as sorbent for H2S removal although in reducing environments ZnO can be reduced to Zn and this might evaporate. Table 6 presents the residual concentrations of H2S, COS an Zn as a function of sorbent temperature. Above 400 °C, H2S concentration is very close to the target value of 1 ppmv. Despite thermodynamic equilibrium results suggest an operating temperature up to around 400 °C, Tamhankar et al. showed that experimentally H2S concentration can be lowered below the expected equilibrium concentration due to the formation of a sulfide layer on the surface with free energy lower than the bulk sulphide [37]. The concentration of COS drops significantly below the ppmv value, and it is even lower than the ppbv value below 400 °C. The concentration of gaseous Zn results 1.61 ppbv at 450 °C and below the ppbv threshold at lower temperatures. Therefore, in this work the evaporation of zinc was not considered a threat for the system.

Although HCl concentration could be decreased well below the target value at temperatures higher than 400 °C, ZnO should not be operated above this temperature. Therefore, the GCU operating temperature was fixed at 400 °C. The calculation also showed that...
HCl reacts with ZnO forming ZnCl₂. This does not affect the removal of H₂S and COS but the equilibrium concentration of gaseous ZnCl₂ is between 2.5 and 5.3 ppmv in the temperature interval under analysis. This compound might damage the SOFC. Moreover, at 300 and 325 °C liquid ZnCl₂ might form. Conversely, S compounds do not react with K₂CO₃. Therefore, HCl removal should be carried out before that of H₂S.

| Table 2 | Model input parameters. |
| --- | --- |
| **Inlet Flows** | | |
| Biosyngas temperature (SLIP-GAS) | 440 | [ °C] |
| Steam temperature (SLIP-WT) | 260 | [ °C] |
| SOFC air temperature (SOFC-AIR) | 25 | [ °C] |
| **Gas Cleaning Unit (GCU)** | | |
| Outlet syngas temperature | 395 | [ °C] |
| Heat losses | 400 | [W] |
| **Tar reformer (TAR-REF)** | | |
| Reactor temperature | 740 | [ °C] |
| Heat losses | 160 | [W] |
| **Heat exchanger (ANODE-HX)** | | |
| Heat losses | 160 | [W] |
| **CPOX unit** | | |
| Heat losses | 160 | [W] |
| **SOFC unit** | | |
| Fuel utilization (based on SLIP-GAS) | 0.8 | | |
| Cell active area | 127 | [cm²] |
| Area specific resistance | 0.35 | [Ω cm²] |
| Efficiencies | | |
| Outflows temperature | 830 | [ °C] |
| SOFC reaction temperature | 830 | [ °C] |
| DC power | 6000 | [W] |
| Heat losses | 900 | [W] |
| **Air heat exchanger (AIR-HX)** | | |
| Air outlet temperature | 700 | [ °C] |
| Flue gas outlet temperature | 180 | [ °C] |
| Heat losses | 160 | [W] |
| **Power electronics (W-CDTNR)** | | |
| Electric efficiency | 95 | [%] |
| **After burner (AFTBURN)** | | |
| Heat losses | 220 | [W] |
| **Heat recovery unit (HT-RECOV)** | | |
| Heat losses | 160 | [W] |
| **SOFC compressor (SOFC-CMP)** | | |
| Isentropic efficiency | 70 | [%] |
| Electric efficiency | 95 | [%] |
| **Others** | | |
| Pressure drop in components | 1 | [%] |

| Table 3 | Gas composition after slip stream humidification. |
| --- | --- |
| Compound | Woodchips [wt% w.b.] | Straw pellets [wt% w.b.] |
| CO | 12.01 | 10.10 |
| H₂O | 33.33 | 33.53 |
| CO₂ | 15.15 | 15.53 |
| H₂ | 0.92 | 0.80 |
| CH₄ | 0.67 | 0.68 |
| C₂H₄ | 0.62 | 0.62 |
| gravimetric tar | 4.80 | 5.80 |
| formaldehyde | 0.31 | 0.37 |
| propene | 0.43 | 0.52 |
| acetic acid | 1.72 | 2.08 |
| methanol | 0.12 | 0.15 |
| ethanol | 0.74 | 0.89 |
| N₂ | 29.18 | 28.93 |

| Table 4 | Equilibrium sulphur and chlorine compounds concentrations at 400 °C. |
| --- | --- |
| Species | molar fraction |
| **Woodchips** | | |
| HCl | 8.1E-06 |
| H₂S | 2.4E-05 |
| COS | 3.5E-08 |
| **Straw pellets** | | |
| HCl | 8.7E-04 |
| H₂S | 2.1E-04 |
| COS | 3.2E-07 |
4.2. Tar sampling and analysis

Gravimetric analysis indicated a tar amount of 120–130 g Nm$^{-3}$ dry biosyngas for woodchips and 190 g Nm$^{-3}$ for straw pellets. The results of the GC-MS analysis are reported in Fig. 3, where only the compounds with a concentration higher than 1 g Nm$^{-3}$ dry biosyngas are presented. Some of the GC-MS measured compounds, such as acetic acid and methanol, were already included in biosyngas composition since they were measured with FT-IR, as explained in the methodology. The sum of all the species analysed was equal to 100–110 g Nm$^{-3}$ of biosyngas for woodchips and 245 g Nm$^{-3}$ for straw pellets. These amounts are different from the one measured with the gravimetric analysis. Gravimetric tar from straw pellets was less than the GC-MS measured tar, while for woodchips the opposite was observed (Fig. 4). This might be due to the lower moisture content of the fuel, 8 wt% as compared to 30 wt% for woodchips. Less water in the fuel resulted in higher fuel bed temperature in the gasifier. As a consequence, more pyrolytic lignin was converted into volatile organic compounds, which are detectable with GC-MS but not with gravimetric analysis.

Table 7 presents the gravimetric tar elemental composition. The large amount of oxygen indicates that the mixture was mostly composed of oxygenated compounds, in agreement with what reported in literature for updraft gasifiers and with the results of the GC-MS analysis [30,39].

The weight percentages were converted into number of moles and an empirical formula for gravimetric tar was calculated. The presence of nitrogen was considered negligible. Gravimetric tar was represented with the molecule C$_4$H$_8$O$_2$ for woodchips and C$_4$H$_8$O$_2$ for straw pellets. Three compounds were selected and their ration was calculated to match the empirical formula. The compounds selected to represent gravimetric tar are 2-methoxyphenol, hydroxyaceton and hydroxyacetic acid. While the first two compounds were measured with the GC-MS analysis, the last compound was selected to close the elemental mass balance from a list of primary tar reported in Ref. [30]. Table 8 reports the weight percentages of the selected tar compounds in the humidified biosyngas.

The compounds selected are representative of both the carbohydrate-derived and lignin-derived part of the feedstock [40,41]. The three compounds have boiling point well above 60 °C, that is the temperature used in the gravimetric analysis. Moreover, their vapour pressure is significantly lower than that of isopropanol and acetic acid, as visible from the values reported in Table 9. Therefore, they were assumed to remain in the gravimetric sample. Analysis with GC-MS of the gravimetric residue dissolved in isopropanol could give an indication of the accuracy of this

| Temperature [°C] | HCl molar fraction | H2S molar fraction | COS molar fraction | Zn molar fraction |
|-----------------|--------------------|--------------------|-------------------|------------------|
| 350             | 2.2E-07            | below ppb          | below ppb         |                  |
| 375             | 3.8E-07            | below ppb          | below ppb         |                  |
| 400             | 6.2E-07            | below ppb          | below ppb         |                  |
| 425             | 9.6E-07            | 1.9E-09            | below ppb         |                  |
| 450             | 1.4E-06            | 3.6E-09            | 1.6E-09           |                  |

Table 5

| Temperature [°C] | H2S molar fraction | COS molar fraction | Zn molar fraction |
|-----------------|--------------------|--------------------|------------------|
| 350             | 2.2E-07            | below ppb          | below ppb         |                  |
| 375             | 3.8E-07            | below ppb          | below ppb         |                  |
| 400             | 6.2E-07            | below ppb          | below ppb         |                  |
| 425             | 9.6E-07            | 1.9E-09            | below ppb         |                  |
| 450             | 1.4E-06            | 3.6E-09            | 1.6E-09           |                  |

Table 6

The weight percentages were converted into number of moles and an empirical formula for gravimetric tar was calculated. The presence of nitrogen was considered negligible. Gravimetric tar was represented with the molecule C$_4$H$_8$O$_2$ for woodchips and C$_4$H$_8$O$_2$ for straw pellets. Three compounds were selected and their ration was calculated to match the empirical formula. The compounds selected to represent gravimetric tar are 2-methoxyphenol, hydroxyaceton and hydroxyacetic acid. While the first two compounds were measured with the GC-MS analysis, the last compound was selected to close the elemental mass balance from a list of primary tar reported in Ref. [30]. Table 8 reports the weight percentages of the selected tar compounds in the humidified biosyngas.

The compounds selected are representative of both the carbohydrate-derived and lignin-derived part of the feedstock [40,41]. The three compounds have boiling point well above 60 °C, that is the temperature used in the gravimetric analysis. Moreover, their vapour pressure is significantly lower than that of isopropanol and acetic acid, as visible from the values reported in Table 9. Therefore, they were assumed to remain in the gravimetric sample. Analysis with GC-MS of the gravimetric residue dissolved in isopropanol could give an indication of the accuracy of this

![Fig. 3. GC-MS analysis results for woodchips (dashed) and straw pellets (solid).](image-url)
assumption.

4.3. System modelling

The first results of the modelling activities are the SOFC operating conditions, as illustrated in Fig. 5. There is no significant difference in operating conditions between the two fuels in analysis. Similarly, the option for internal or external reforming (case D,C vs case A,B) has negligible impact on the operating conditions of the fuel cell. Conversely, substantial differences are visible depending on the biosyngas heating up strategy adopted (case A,D vs case B,C). Using the afterburner off gases results in higher reversible voltage and cell voltage. This is due to the higher H₂ concentration in the biosyngas reaching the anode.

The model was then used to calculate the flow rates necessary to generate 6 kW in the SOFC keeping the fuel utilization equal to 0.8. Table 10 illustrates the mass flow rates of biosyngas, steam, cathode air, and CPOX air. When part of the biosyngas is combusted in the CPOX unit (cases B and C), more biosyngas is necessary to generate the targeted 6 kW. As a consequence, also the flow rate of water to be added to the slip stream to avoid carbon deposition increases. In all the cases analysed, the biosyngas flow rate from straw pellets is lower than the one from woodchips. This result shows the importance of using the energy content of light condensable and volatile organic compounds, and gravimetric tar. In fact, straw pellets-derived biosyngas has a lower content of CO and almost equal content of H₂ and CH₄, but it has a larger content of these compounds. The data also show clearly the reduction in SOFC cooling needs due to direct internal tar reforming (cases C and D). In case of woodchips-derived biosyngas, the cathode air decreases from approximately 87 kg h⁻¹ to 61 kg h⁻¹ and in case of straw pellets-derived biosyngas, due to the larger tar content, the SOFC air reduces even further from around 87 kg h⁻¹ to 57 kg h⁻¹.

The model was then used to compare the energy performances of the different system configurations. Table 11 shows the results of the system energy analysis. The required energy input varies between roughly 14 and 17 kW depending on the biosyngas heating-up strategy adopted, in accordance with the previously discussed larger biosyngas flowrate necessary to generate the target 6 kW in the SOFC when the CPOX unit is used. In the system, between 4 and 6 kW are lost. The two cases with external reformer are the ones with the largest total energy losses, equal to 5.3 kW and 5.7 kW when the afterburner off gases and the CPOX unit are used to heat up the biosyngas, respectively. The reason for this behaviour can be

Table 7
Gravimetric tar elemental composition.

| Element | Woodchips [wt% w.b.] | Straw pellets [wt% w.b.] |
|---------|----------------------|--------------------------|
| C       | 58.70                | 56.90                    |
| H       | 6.30                 | 6.50                     |
| O       | 34.90                | 36.60                    |
| N       | 0.16                 | 0.10                     |

Table 8
Amount of the selected representative tar in the humidified biosyngas.

| Compound                | Woodchips [wt% w.b.] | Straw pellets [wt% w.b.] |
|-------------------------|-----------------------|--------------------------|
| 2-Methoxyphenol         | C₇H₇O₂                 | 3.51                     |
| hydroxyacetic acid      | C₄H₇O₂                 | 1.07                     |
| Hydroxyaceton           | C₃H₆O₂                 | 0.22                     |

Table 9
Boiling temperature and vapour pressure at 25 °C of the selected compounds.

| Compound         | Boiling Temperature [°C] | Vapour pressure [mm Hg] |
|------------------|--------------------------|-------------------------|
| Isopropanol      | 83                       | 45.4                    |
| Acetic acid      | 118                      | 15.7                    |
| Hydroxyaceton    | 145                      | 2.95                    |
| 2-methoxyphenol  | 205                      | 0.103                   |
| Hydroxyacetic acid| 100                     | 0.0202                  |

Fig. 4. Comparison of gravimetric (dashed) and GC-MS (solid) analysis.
ascribed to the higher chimney losses that are higher when external reformer is used since a larger cathode air flow rate is necessary to maintain the SOFC operating temperature. The energy losses at the chimney, equal to around 3 kW in case of external reformer and 2 kW in the configuration with direct internal reforming, account for the largest losses, as visible in Fig. 6, where the share of the losses in the configuration with direct internal reforming and afterburner off gases utilization (case D) is presented for woodchips-derived biosyngas. On top of decreasing the chimney energy losses, the lower cathode flow rate achievable with direct internal reforming results also in lower energy required by the SOFC compressor, that decreases from 0.6 to 0.4 kW with both fuels and in both CPOX and afterburner off gases configurations. As a consequence, the net energy output also increases marginally. Using the CPOX unit results in slightly higher losses due to the presence of an additional component, thus an increased heat loss; moreover, the higher biosyngas flow rate required to produce the

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**Table 10** Biomass, steam, cathode air and, CPOX air mass flow rates.

| Case | Woodchips | Straw pellets |
|------|-----------|--------------|
|      | A | B | C | D | A | B | C | D |
| Max flow rate [kg h⁻¹] | SLIP-GAS | 8.46 | 10.42 | 10.45 | 8.48 | 7.38 | 9.08 | 9.11 | 7.39 |
| SLIP-WT | 1.02 | 1.25 | 1.25 | 1.02 | 1.84 | 2.27 | 2.28 | 1.85 |
| Int. air (CPOX) | – | 2.92 | 2.90 | – | – | 2.88 | 2.89 | – |
| SOFC-Air | 86.85 | 86.66 | 60.9 | 61.29 | 86.82 | 86.70 | 56.49 | 56.89 |

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**Table 11** Results of system energy analysis.

|          | Woodchips | Straw pellets |
|----------|-----------|--------------|
|          | A | B | C | D | A | B | C | D |
| Energy input [kW] | SLIP-GAS | 13.75 | 17 | 17.25 | 13.82 | 13.46 | 16.57 | 16.62 | 13.49 |
| SLIP-WT | 0.13 | 0.16 | 0.16 | 0.13 | 0.23 | 0.29 | 0.29 | 0.23 |
| Total | 13.88 | 17.16 | 17.31 | 13.95 | 13.69 | 16.86 | 16.91 | 13.72 |
| Energy losses [kW] | SOFC | 0.9 | 0.9 | 0.9 | 0.9 | 0.9 | 0.9 | 0.9 |
| Other reactors | 1.1 | 1.26 | 1.1 | 1.1 | 1.1 | 1.26 | 1.1 |
| DC/AC | 0.3 | 0.3 | 0.3 | 0.3 | 0.3 | 0.3 | 0.3 |
| Chimney | 3.04 | 3.23 | 2.3 | 2.3 | 3.03 | 3.21 | 2.28 | 2.12 |
| Total | 5.34 | 5.69 | 4.6 | 4.6 | 5.33 | 5.67 | 4.58 | 4.12 |
| Energy transmitted [kW] | SOFC-CMP | 0.55 | 0.58 | 0.41 | 0.41 | 0.55 | 0.58 | 0.41 | 0.38 |
| W-NET | 5.15 | 5.12 | 5.29 | 5.29 | 5.15 | 5.12 | 5.29 | 5.32 |
| HT-RECOVER | 3.69 | 6.26 | 7.49 | 4.36 | 3.48 | 6.38 | 7.35 | 4.06 |
| Total | 8.84 | 11.77 | 12.61 | 9.65 | 8.63 | 11.5 | 12.64 | 9.38 |
target 6 kW and the additional presence of 2.9 kg h⁻¹ air for the CPOX operation further increase the chimney losses. The choice between using the afterburner off gases or the CPOX unit significantly affects also the amount of heat available in the recovery section. In case of woodchips as fuel, the heat recovered decreases when using the afterburner off gases from 6.3 kW to 3.7 kW with external reforming, and from 7.5 kW to 4.4 kW with direct internal reforming. The same trend holds true for pellet as fuel. Using the afterburner off gases to heat up the biosyngas and to sustain the reforming reactions in the external reformer (case A) results in the lowest amount of heat available, equal to 3.5 kW with straw pellets-derived biosyngas. Although to a minor extent, also the reforming strategy affects the heat that can be recovered, that increases from 3.7 kW to 4.4 kW for woodchips and from 3.5 kW to 4.0 kW for straw pellets in the configuration with after burner off gases utilization.

Fig. 7 shows the electrical, thermal, and total efficiency achievable with woodchips- and straw pellets-derived biosyngas in the different system configurations. The equations used to calculate the efficiencies can be found in Appendix A.2. The highest overall efficiency, that is 74% with woodchips and 75% with straw pellets, is achieved with direct internal tar reforming and using the CPOX unit to increase the biosyngas temperature before the SOFC. Using the CPOX unit results in higher overall efficiencies due to an increase in the thermal efficiency. However, it significantly lowers the system electrical efficiency to around 30% as compared to roughly 37% achievable when the afterburner off gases are used to heat up the biosyngas. Fuel cells are electrochemical devices and therefore to achieve high voltages and efficiencies, it is important to keep high the chemical energy of the fuel before it enters the fuel cell. The lower cathode air required due to direct internal reforming slightly reduces the electricity consumption by the compressor and the system electrical efficiency increases from 37.1% to 38.0% with woodchips as fuel and from 37.6% to 38.8% with straw pellets in the configurations with the afterburner off gases to heat up the biosyngas. In the configuration with the CPOX unit, direct internal reforming increases the system electrical efficiency from 30.4% to 31.3% with pellets and 29.9%–30.6% with woodchips. Direct internal reforming also increases the system thermal efficiency thanks to the higher amount of heat available in the recovery section and the lower heat depleted through the chimney thanks to the decreased cathode flow.

The results obtained in the energy analysis are also presented in the form of energy flows. Fig. 8 illustrates the two cases with external reformer and woodchips as fuel. Focusing on the amount of chemical energy and thermo-mechanical energy, it can be seen that for case A, there is an increase in chemical energy of the fuel flow that exits the tar reformer. The reforming reactions produce additional chemical energy with the help of the thermo-mechanical energy provided by the afterburner off gases. On the other hand, when the CPOX unit is employed as in case B, a part of the chemical energy is converted into thermo-mechanical energy when the gas passes through the CPOX unit. Considering that the slip stream extraction is used for electricity production, it might appear reasonable to design the system to have the highest electrical efficiency, that is using the afterburner off gases to heat up the biosyngas.

The efficiency results obtained from the system modelling can be compared with the results achieved with the real system during the Horizon2020 project "FlexiFuel-SOFC". These results are not used as validation of the model, but rather as additional information on the performance that can be actually achieved with an integrated biomass gasifier SOFC CHP system. The configuration of the real system is slightly different from the one adopted in the model since, after having gained experience with a first pilot plant, a second improved pilot plant was built. Moreover, as initially mentioned, the model does not include the gasifier, but it starts from the slip stream and it ends with the boiler, where only the afterburner off gases are used to generate heat. Therefore, the energy input in the model is the biosyngas extracted in the slip stream, while in the real system it is the solid biomass fed to the gasifier. Furthermore, in the real system the biosyngas that is not extracted with the slip stream is used to generate additional heat. Last, despite the successful operation of SOFC with tar-containing biosyngas, since there is not yet general agreement on the fate of tar in the anode chamber, in the system tar was converted before reaching the SOFC [42]. A total efficiency of 80% was achieved with the real system and this can be further increased to 91% by adopting a condenser in the heat recovery section. The SOFC used the biosyngas to generate electricity with a gross DC efficiency higher than 42%, and the heat recovered from the SOFC exhaust accounted for 21% of the total system energy output. The project results have been

![Fig. 7. Electrical, thermal, and total efficiency with (top) woodchips- and (bottom) straw pellets-derived biosyngas.](image-url)
Fig. 8. Energy flows of the two system configurations with external reformer and woodchips as fuel.
presented during the final project workshop held during the European Biomass Conference and Exhibition (EUBCE) 2019 in Lisbon. More details can be found on the project website flexifuelsocfu.eu, where the presentations given in the workshop are available for download [43].

The model developed in this study will be further improved in our future work. The gasification step will be included and the system configuration will be revised to represent the final “Flexifuel-SOFC” system design. Moreover, the values used in terms of pressure drops and heat losses will be updated in light of the measurement performed with the real system. The model will be successively validated thanks to the results of the extensive experimental campaign carried out within the project.

5. Conclusion

This work aimed to improve the competitiveness of an integrated biomass updraft gasifier SOFC micro CHP system. This was done by increasing system efficiency and reducing its complexity via direct internal reforming of hydrocarbons. We used a comprehensive approach combining system modelling, thermodynamic equilibrium calculations and experimental data to evaluate the efficiency gain obtainable, identify an appropriate gas cleaning configuration, and compare two strategies to heat up the biosyngas before feeding it to the SOFC: catalytic partial oxidation and afterburner off gases utilization.

The GCU has to operate at temperatures lower than 400 °C to decrease H2S and HCl below SOFC tolerance limit using ZnO and K2CO3 as sorbents. However, HCl removal should be carried out before that of H2S to avoid reactions between ZnO and HCl leading to the formation of ZnCl2. Gravimetric analysis indicated a tar amount of 120–130 g Nm−3 dry biosyngas for woodchips and 190 g Nm−3 for straw pellets. Differently, the sum of all the species analysed with GC-MS analysis was equal to 100–110 g Nm−3 dry biosyngas for woodchips and 245 g Nm−3 for straw pellets. The majority of the tar measured with GC-MS analysis belongs to the primary tar group, and 2-methoxyphenol, hydroxyacetic acid and hydroxyacetone were selected as representative compounds.

Direct internal tar reforming reduces the SOFC cooling needs. In case of woodchips-derived biosyngas, the cathode air decreases from approximately 87 kg h−1 to 61 kg h−1 and from around 87 kg h−1 to 57 kg h−1 in case of straw pellets-derived biosyngas. The lower cathode air required due to direct internal reforming reduces the electricity consumption by the compressor and the system electrical efficiency increases from 37.1% to 38.0% with woodchips as fuel and from 37.6% to 38.8% with straw pellets in the configuration with the afterburner off gases used to heat up the biosyngas. In the configuration with the CPOX unit, direct internal reforming increases the system electrical efficiency from 30.4% to 31.3% with pellets and 29.9%–30.6% with woodchips. When tar is reformed internally also the heat that can be recovered increases.

Using the CPOX unit results in higher overall efficiencies due to an increase in the thermal efficiency. The heat recovered increases when using the CPOX unit. However, it significantly lowers the system electrical efficiency to around 30% as compared to roughly 38% achievable when the afterburner off gases are used to heat up the biosyngas. Since the system is used for electricity production, it is reasonable to adopt the configuration with the highest electrical efficiency, that is with after burner off gases utilization.

A simplified system design can be obtained with direct internal reforming, which might also have a significant impact on both capital and operation costs of the system. A techno-economic analysis can be used to evaluate the advantages at system level. From an energy point of view, direct internal reforming brings an increase in electrical efficiency of around 1% point and in thermal efficiency of even 5–6% points. The choice of direct internal tar reforming seems therefore an advantageous strategy to improve the competitiveness of an integrated biomass updraft gasifier SOFC micro CHP system.

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.

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Appendix A. Thermodynamic approach

A.1 SOFC modelling

Aspen plus is not provided with an SOFC model. Therefore, a combination of various blocks and a fortran-based code written in a calculator block is needed for SOFC modelling and to perform the electrochemical calculations.

The reversible voltage (Erev ) is calculated as:

\[
E_{\text{rev}} = \frac{R \cdot T_{\text{SOFC}}}{4 \cdot F} \cdot \ln \left( \frac{P_{\text{o2, cathode}}}{P_{\text{o2, anode}}} \right)
\]

where \( R \) is the ideal gas constant (J/kg mol), \( T_{\text{SOFC}} \) is the operating temperature of the fuel cell (K), \( p \) is the partial pressure of the oxygen in the anode and cathode sides (Pa) and \( F \) is the Faraday’s constant (C/mol) [44].

The cell voltage (Vcell) is determined as:

\[
V_{\text{cell}} = E_{\text{rev}} - j \cdot ASR
\]

Being \( j \) is the current density (A/m^2) and ASR is the area-specific resistance (Ω m^2) [45].

The biosyngas flow rate is calculated by determining the equivalent H2 content of the anode inlet flow. The equivalent H2 content is given by the content of hydrogen available after complete reforming and water gas shift of the hydrocarbons.

A.2 Efficiency calculations

The electric efficiency is calculated as

\[
\eta_{\text{electric}} = \frac{\dot{W}_{\text{SOFC}} - \dot{W}_{\text{DC/AC}} - \dot{W}_{\text{SOFC-CMP}}}{m_{\text{SLIP-ST}} \cdot (LHV + 2\Delta h) + m_{\text{SLIP-WT}} \cdot 2\Delta h}
\]

The energy of biosyngas (SLIP-ST) is given by the lower heating value (LHV in kJ/mol) added to the enthalpy variation (\( \Delta h \) in kJ/kg) from 25 °C to 440 °C multiplied by its mass flow (kg/s). To remark that, from the enthalpy variation calculation for both biosyngas and steam (SLIP-WT), the latent heat of water is subtracted. The net electricity \( \dot{W}_{\text{NET}} \) in kW is given by the SOFC electricity production \( \dot{W}_{\text{SOFC}} \)
subtracted by the electric losses in the power electronics ($W_{DC/AC}$) and the electricity supplied to the compressor ($W_{SOFC-CMP}$).

The thermal efficiency is calculated as

$$\eta_{thermal} = \frac{Q_{HT \_RECOV}}{m_{SLIP \_ST} \cdot (LHV + D\eta) + m_{SLIP \_WT} \cdot D\eta}$$

$$\eta_{thermal} = \frac{Q_{HT \_RECOV}}{m_{SLIP \_ST} \cdot (LHV + D\eta) + m_{SLIP \_WT} \cdot D\eta}$$

($kW$) represents the heat transferred to the domestic hot water in the heat recovery section [46].

The total efficiency ($\eta_{total}$) of the system is calculated as

$$\eta_{total} = \eta_{electric} + \eta_{thermal}$$

Credit author statement

A. Cavalli: Conceptualization, Methodology, Experiments, Modelling, Visualisation, Writing – review & editing. A. Fernandes: Conceptualisation, Modelling, Writing, Reviewing, Visualization, P. V. Aravind: Supervising, Reviewing.

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