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Study of CO2 recovery in a carbonate fuel cell tri-generation plant

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HIGHLIGHTS
- Separation of CO2 in a biogas plant that co-produces electricity, hydrogen, and heat.
- The ability of MCFC is to concentrate CO2 in the anode exhaust stream.
- Three cathode inlet configurations are considered.
- Results illustrate a high compatibility between hydrogen co-production and CO2 recovery.
- A series configuration of MCFC technology coupled with an ICE achieves outstanding carbon recovery (exceeding 90%).

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ABSTRACT
The possibility of separating and recovering CO2 in a biogas plant that co-produces electricity, hydrogen, and heat is investigated. Exploiting the ability of a molten carbonate fuel cell (MCFC) to concentrate CO2 in the anode exhaust stream reduces the energy consumption and complexity of CO2 separation techniques that would otherwise be required to remove dilute CO2 from combustion exhaust streams. Three potential CO2 concentrating configurations are numerically simulated to evaluate potential CO2 recovery rates: 1) anode oxidation and partial CO2 recirculation, 2) integration with exhaust from an internal combustion engine, and 3) series connection of molten carbonate cathodes initially fed with internal combustion engine (ICE) exhaust. Physical models have been calibrated with data acquired from an operating MCFC tri-generating plant. Results illustrate a high compatibility between hydrogen co-production and CO2 recovery with series connection of molten carbonate systems offering the best results for efficient CO2 recovery. In this case the carbon capture ratio (CCR) exceeds 73% for two systems in series and 90% for 3 MCFC in series. This remarkably high carbon recovery is possible with 1.4 MWe delivered by the ICE system and 0.9 MWe and about 350 kg day−1 of H2 delivered by the three MCFC.

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1. Introduction
The global impacts of CO2 emissions from power generation have been well established [1]. Industrialized nations have made strong commitments for the reduction of CO2 and other greenhouse gases emissions [2]. This presents a considerable technical challenge and significant driver for change to the power industry. Leading solutions include the introduction of large amounts of renewable power, more efficient systems for electricity production, more efficient transmission and use of electricity, and carbon capture and sequestration from fossil fuel plants. Most projections for future power generation include substantial contributions from fossil fuel combustion [3] to complement the cost, control and intermittency features of renewable power generation. Carbon separation in traditional fossil fuel power plants can be achieved either upstream or downstream of the combustion zone using a variety of technologies including cryogenics, air separation, and oxy-combustion [4], [5], [6]. Moreover, in recent years the opportunities for CO2 recovery have grown considerably. Pure captured CO2 is indeed a potential valuable feedstock for the synthesis of chemicals and synthetic fuels. Utilization rather than geological sequestration is also related to the poly-generation approach, i.e., energy systems featuring the co-production of several products (heat, electric power and chemicals).
In this paper, a poly-generation system based on molten carbonate fuel cell is first studied, and new structures of the system are analyzed in order to provide a carbon recovery added effect. The transportation sector is searching for alternatives to traditional liquid hydrocarbon fuels by considering electricity, natural gas, biofuels and hydrogen, amongst other alternative fuels. Co-production of electricity and hydrogen with a molten carbonate fuel cell presents a unique opportunity for synergistic co-production with efficient carbon capture. High temperature fuel cells, especially molten carbonate (MCFC) and solid oxide fuel cells (SOFC), have received substantial attention during the past decades and have recently begun to achieve commercial success [7], [8]. Both MCFC and SOFC technologies can operate on natural gas and biogas to achieve higher fuel-to-electric (FTE) efficiencies and near zero criteria pollutant emissions (e.g. NOx, SOx, particulate) compared to traditional engines. Efficiency is further increased through the concept of “tri-generation”, in which electrical power, thermal power and hydrogen are co-produced from a single fuel stream [9]. Tri-generation utilizes waste heat and chemical recovery concepts to produce a viable fuel that can be used for zero emission transportation purposes, as promoted by the state of California since the year 1990 [10]. With natural gas as a feedstock, CO2 emissions can be lowered between 10 and 43.6% with a high temperature fuel cell, depending upon the electric power generating system considered in the comparison, while with renewable biogas feedstock the emissions can be reduced near to zero [11].

With this in mind, the current work investigates a further reduction of greenhouse gas emissions adding the recovery of CO2 from a tri-generation MCFC plant. The energy intensity of CO2 recovery is reduced when exploiting the carbonate ion transfer mechanism of molten carbonate fuel cells (Fig. 1). MCFC consume CO2 in the cathode compartment and produce CO2 in the anode compartment where it is only mixed with a small amount of fuel and product constituents (e.g., H2, CO, water). When this anode exhaust stream is fully converted the CO2 is only mixed with water allowing simple and inexpensive pure CO2 recovery by cooling and water condensation.

Subsequent to the anode the remaining hydrogen content of the anode exhaust stream is typically oxidized in a combustor. To eliminate the introduction of nitrogen and excess oxygen the current design utilizes nearly stoichiometric oxy-combustion with oxygen supplied from an air separation unit. The resulting products of combustion, H2O and CO2, are readily separated with considerably less energy input than cryogenic CO2 distillation. The plant configurations outlined in section 5 are an extension of the existing MCFC tri-generation plant located at the Orange County Sanitation District [12]. A molten carbonate fuel cell is fed with biogas produced on-site from the anaerobic digestion of sludge available in the wastewater treatment plant. This study outlines the feasibility of applying carbon sequestration to an already existing and operating MCFC tri-generation plant, with the aim of exploring synergies and limitations arising from the integration of a CO2 recovery unit in a tri-generation plant co-producing electrical power, heat and hydrogen.

The CO2 recovered could be recycled since its purity is sufficiently high. For example it could be sold as a commercial product in the food or chemical industry. Another possible use is the conversion to fuel (e.g., in a photo-bio-reactor with the aid of nutrients (phosphates and nitrates) the added CO2 stream can be converted to algae and oxygen). A third option is the transportation in pipeline to suitable sites for CO2 sequestration. At present, a large share of CCS occurs through the injection in geological deposits previously used for the extraction of oil or gas. The combination of these techniques with a tri-generation system operated on biogas (such as the one at OCSD) would enable carbon-negative production of electricity and fuel (with the CCS process), since the carbon of the ADG was previously contained in the sewage (before being converted into biogas), which ultimately comes from plants that removed CO2 from the atmosphere.

### 2. Background

Molten carbonate fuel cells utilize carbonate ions as the charge carriers in the electrochemical conversion of chemical energy into electrical energy. Operation between 550 and 650 °C is necessary in order to maintain a sufficient electrolyte conductivity and chemical reactivity without the use of noble metal catalysts. The

---

**Nomenclature**

| Symbol | Description |
|--------|-------------|
| ASU    | Air separation unit |
| α      | charge transfer coefficient |
| CCR    | Carbon capture ratio |
| CCS    | Carbon capture and separation |
| CHP    | Combined heat and power |
| F      | Faraday constant [96485 C mol⁻¹] |
| HSU    | Hydrogen separation unit |
| I      | Current [A] |
| ICE    | Internal combustion engine |
| LHVCH4 | Biogas lower heating value [800 MJ kmol⁻¹] |
| MCFC   | Molten carbonate fuel cell |
| NFCRC  | National Fuel Cell Research Center |
| $N_i$  | Molar flow of the species “i” [kmol s⁻¹] |
| $\eta$ | Electrical efficiency |
| OCSD   | Orange County Sanitation District |
| $P_{el}$ | Electric power [kW] |
| $P_{el, ICE}$ | ICE equivalent electric power [kW] |
| PSA    | Pressure swing adsorption |
| STC    | Steam to carbon ratio |
| T      | Temperature |
| $U_f$  | Fuel utilization factor |
| $U_{CO2}$ | CO2 utilization factor |
| V      | Voltage [V] |
| WGS    | Water gas shift |
| $X_i$  | Molar concentration of the species “i” |
| $\gamma_{CH4}$ | Biogas composition — methane molar concentration |

---

**Fig. 1.** Molten carbonate fuel cell.
temperature is well-suited for integration with hydrocarbon reforming in a fuel processor. The typically endothermic reactions which convert common fuels such as methane, biogas or ethanol into a hydrogen-rich feedstock is able to provide a substantial portion of the stack cooling requirement. The reforming process can occur directly within the anode gas channels and/or in separate in-stack plate reformers, as is the case for the current MCFC system.

Use of a MCFC as a carbon dioxide concentrator was investigated experimentally by Sugiura et al. [13], where it is showed that the experimental values for the CO₂ removal rate from cathode to anode performed in a MCFC almost matched theoretical calculations. One key conclusion from this work is that the ratio between the partial pressures of oxygen and carbon dioxide directly affects the performance. Campanari et al. [14] investigated the possibility of separating CO₂ from NG-fed combined cycles integrated with MCFC operating in the exhaust. Results identified the potential for an 80% reduction in CO₂ with negligible impact on electrical efficiency. This reduction was achieved with a MCFC sized to contribute only 20% of the net power generation. A similar study considered cryogenic separation of CO₂ from combined cycles integrated with MCFC [15] by analyzing the possibility of recirculating a portion of the carbon dioxide recovered in the cathode, thereby controlling the utilization of CO₂. This approach avoided the separation via oxy-fuel combustion presented in Campanari et al.

A subsequent paper from the same research group [16] compared three configurations for integrating a MCFC with an internal combustion engine. Carbon emissions from the combined system are dramatically reduced. Separate research has demonstrated this type of plant is, at the moment, un-profitable, but future cost reductions and incentives for fuel cells and avoided carbon emissions make this an attractive technology [17]. The economic analysis considered a small MCFC integrated with a CHP plant capable of separating 63% of the inlet carbon quantity at an overall fuel-to-electric efficiency of 35%. Further improvements in performance may improve the viability of MCFC CO₂ capture as much as potential cost reduction.

### 3. Model description

#### 3.1. General overview on the model

The integration of a MCFC, hydrogen separation and CO₂ separation results in a highly coupled system requiring simulation with detailed physical models of the various sub-systems. This work extends the spatially resolved MCFC system modeling capability of the NFRCC and is comprised of multiple physical models of sub-system components that can be found in the literature [18], [19], [20].

The primary component model is that of the MCFC stack which has been designed and calibrated to represent the physics of the “FuelCell Energy Direct Fuel Cell” (DFC) design. The model permits variable flow geometries to represent other possible stack configurations including external reforming, co-flow, cross-flow and counter-flow manifolding.

System parameters that were calibrated for the bio-gas fed DFC system include the following:

- Active cell area and geometry (channel heights, lengths, and plate thicknesses)
- Fuel utilization (Uᵣ)
- Ohmic and activation polarizations as a function of temperature
- Bio-gas composition
- Steam-to-carbon ratio (STC)

- Air inlet temperature and internal heating

The spatially resolved model output includes principal quantities such as local temperature and species concentrations in both anode and cathode, bulk channel flow rates, and resolved local current density. The balance of plant components include two water gas shift reactors, five heat exchangers, an oxidizer, two water condensers, an evaporator, an air blower, and six plenum volumes.

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#### 4. CO₂ separation process

The objective of the current system integration is to recover carbon dioxide (and hydrogen) from the anode exhaust stream of the MCFC; typically comprised of CO, H₂O (steam), H₂ and CO₂. The water-gas shift and hydrogen separation occurring upstream of the CO₂ removal both serve to increase the CO₂ concentration of the

#### Table 1

| Parameter            | Value |
|----------------------|-------|
| Pₚₑₜ [kW]            | 300   |
| Manifolds disposition| Cross-flow |
| Length [m]           | 1.1   |
| Width [m]            | 0.8   |
| Area [m²]            | 0.88  |
| Thickness of the cell (without gas channels) [m] | 0.0025 |
| Power density [mW cm⁻²] | 120  |
| Number of cells      | 285   |
| Steam-to-carbon ratio (molar) | 2.5:1 |
| Fuel utilization: Uᵣ | 0.65  |
| Bio-gas composition (molar): YCH₄ | 0.65 CH₄, 0.35 CO₂ |
| Reformer typology    | Indirect internal |
| Reformer position    | Every 10 cells |
The anode effluent. Thus the HSU effectiveness directly impacts the recoverable CO2. The components used to achieve a high purity of carbon dioxide are reported in Fig. 2.

The anode exhaust is first cooled to 350 °C prior to the first water gas shift reactor as the carbon monoxide conversion is favored at low temperatures. A second heat exchanger lowers the temperature to 200 °C prior to the low temperature water gas shift reactor. The combination of the two reactors achieves a high conversion of the CO balancing the faster kinetics occurring at 350 °C and the greater conversion efficiency that occur at 200 °C. A subsequent condenser removes most of the water. The gas entering the hydrogen separation unit contains H2 at near 25% mole fraction.

The hydrogen separation process is an energy-intensive process, and the total amount recovered is directly proportional to the electric power consumption. The technology adopted at the existing OCSD plant is pressure swing adsorption (PSA). This process, common in large scale applications, recovers high purity H2 (99.99%). The PSA functions on the principle that gases can be preferentially adsorbed and desorbed from certain surfaces at different pressures [25]. The system recovers approximately 70% of the available hydrogen. Higher recovery rates require significant compression work, exceeding the benefit of the additional hydrogen recovery, since hydrogen must be recirculated to maintain a minimum concentration at the PSA inlet. The PSA at the OCSD plant operates between 10 and 40 bar with an electric load of 58 kW, met by the nearly 300 kW electric output of the MCFC.

Alternative techniques for hydrogen separation such as membrane separation and electrochemical separation could achieve similar integration with carbon recovery at higher net electrical efficiencies [26].

The off-gas stream (remaining constituents) exiting the PSA is comprised primarily of CO2, H2O, and H2 with trace amounts of CO and CH4. The most effective means of converting the remaining quantities of CH4, CO and H2 without introducing nitrogen is stoichiometric combustion with pure oxygen.

Air separation is a complex and energy-intensive process that could represent a substantial additional parasitic load on the MCFC output. However, the oxygen flow requirement for this particular use is small due to the small amount of hydrogen and carbon monoxide remaining in the gas mixture after the WGS and the HSU. At this scale it is most effective for the high purity oxygen to be purchased from an external producer. Significantly larger plants could consider on-site air separation. The current model includes the energy requirement of air separation (as if it was accomplished on-site), but not that of transportation and delivery, when calculating the system performance characteristics and efficiency. The production of high purity oxygen is assumed to require 0.21 kWh kgO2 [22], which can be expressed also as 0.756 kW (gO2) s. It is readily apparent that a higher recovery rate of hydrogen in the HSU reduces the required oxygen flow and thus the parasitic energy consumption in the ASU.

The final process of carbon recovery is the removal of water in a second condenser. The resulting carbon dioxide purity should exceed 96% by volume, depending upon the water removal capability of the second condenser. The total quantity of CO2 recovered through this process is equal to the carbon present in the fuel minus the quantity vented to the atmosphere for configuration 1 and equal to the fuel plus the quantity of CO3 ions that pass across the electrolyte for configurations 2 and 3 where the cathode carbon dioxide is supplied from the exhaust of an ICE. These configurations will be described in detail in the next section.

5. System description

This study will analyze three configurations for using MCFC to recover CO2. These systems differ substantially in their net electrical efficiency and carbon recovery. The three systems are:

- Recirculation of the CO2 from anode exhaust to cathode
- Separation and capture of internal combustion engine exhaust
- Enhanced CO2 recovery with multiple MCFC stacks in series

5.1. Configuration 1: recirculation of the CO2

In this case the only carbon source is the fuel supplied to the FC, and a portion of the anode exhaust is recirculated into the cathode to supply the necessary amount of carbon dioxide. This is similar to the design of existing commercial MCFC systems. Calculation of the CO2 availability in the cathode is achieved using the Faraday’s law expressed for this application in Equation (1).
\[ N_{\text{CO}_2, \text{cath \ in}} = \frac{I}{2F} U_{\text{CO}_2} \]  

(1)

The current is directly measured, thus control of the carbon dioxide stream entering the cathode is possible with any arbitrary selection of a \( \text{CO}_2 \) utilization factor. Lower \( \text{CO}_2 \) utilization factors improve cell voltage marginally, however unutilized cathode \( \text{CO}_2 \) will escape into the environment with the cathode exhaust. Objectively, limiting the carbon dioxide concentration in the cathode exhaust will maximize the recoverable \( \text{CO}_2 \). Recovery of \( \text{CO}_2 \) from the cathode stream would require energy-intensive methods such as cryogenic separation or PSA.

The carbon recovery potential of different system configurations can be evaluated by establishing a \( \text{CO}_2 \) capture ratio (CCR). Equation (2) defines the CCR for this configuration as the maximum potential \( \text{CO}_2 \) recovery as a proportion of the fuel input of the MCFC. This expression assumes all carbon present in the anode outlet is converted to \( \text{CO}_2 \) and separated after two water gas shift reactors, an oxy-combustor, and a condenser. The quantity “\( 1-\text{CCR} \)” embodies the flow of \( \text{CO}_2 \) released into the atmosphere. Due to inefficiencies and leakage in the HSU, and the condenser, real-world CCR will be only slightly reduced.

\[
\text{CCR} = \frac{N_{\text{CO}_2, \text{cath \ out}} = \frac{N_{\text{Carbon to MCFC}} - N_{\text{CO}_2, \text{cath \ in}} \times (1 - U_{\text{CO}_2})}{N_{\text{Carbon to MCFC}}}}{N_{\text{Carbon to MCFC}}}
= 1 - \left( \frac{1 - U_{\text{CO}_2}}{U_{\text{CO}_2}} \right) \]  

(2)

The pinch analysis methodology was applied for determining the optimal heat integration among the various heat demands/sources within the analyzed plant. The system does not require an external heat source and is capable of recovering thermal power at moderate and low temperatures from the cathode exhaust and the two condensers on the anode stream. The three sources of heat are the outlet streams from the anode (\(-620\) °C), the cathode (\(-650\) °C) and the oxidizer (\(-500\) °C). The temperature difference between cathode and anode outlet is due to manifolds disposition (cross-flow) and cell geometry (rectangular). Those characteristics depend on the MCFC technology adopted by the producer [7]. Cathode flow rate is controlled with a blower in order to maintain a temperature difference of 100 °C between the inlet and the outlet of the cathode compartment. Cathode recirculation is another option for controlling air flow, but is not applied in the case of the DFC design. A detailed plant schematic for configurations 1 and 2 is presented in Fig. 3 (the values of each cornerstone are reported in the Appendix A). The dashed line represents the internal \( \text{CO}_2 \) recirculation of configuration 1. The pressure losses for an ambient pressure MCFC are responsible for the sizable cathode blower parasitic, but are not significant enough to affect the chemical kinetics, so they have been assumed constant in the current study. The baseline operating conditions for the MCFC tri-generation plant are specified in Table 2.

At a carbon dioxide utilization factor of 80%, Equation (2) determines a CCR of 56.4%. A carbon dioxide utilization factor of 80% implies a relatively low voltage which is undesirable. At \( U_{\text{CO}_2} \) values higher than 90% (Fig. 4), the \( \text{CO}_2 \) starvation phenomenon takes place and concentration losses significantly affect the total voltage. In configurations 1 and 2 the maximum \( U_{\text{CO}_2} \) value remains below this threshold, operating at an 80% utilization, in order to avoid issues of reactant starvation in the downstream regions of the
MCFC. This represents a practical operating condition for the MCFC which avoids the precipitous voltage drop at higher CO₂ utilizations. Note that examinations at very low reactant concentrations are only observable with a spatially discretized model, such as the one used in this study, where flow geometry partially mitigates the sudden onset of diffusion loses at high CO₂ utilizations.

Besides the electrochemical losses in the fuel cell, one must also consider the electric power consumed by the air blower and PSA compressors. Considering both the net electricity output and the chemical potential of the recovered hydrogen the co-production efficiency rises to 60.9% of the lower heating value of the biogas fuel. The recovered CO₂ purity is high (0.978) when the second water condenser effectiveness is similarly high.

**5.2. Sensitivity analysis — Configuration 1**

Fig. 5 shows the effects of fuel utilization variation, which can be manipulated by adjusting either fuel flow or current, or some combination thereof. Variation of U₉ illustrates the direct dependence of CO₂ removal on H₂ separation. Increasing fuel utilization reduces the hydrogen recovery, and subsequently CO₂ recovery. Voltage drops, while net electrical efficiency rises. This relative low value is caused mainly by the HSU electrical consumption. Cathode air flow increases in proportion to the reduced voltage in order to maintain a constant internal temperature and temperature gradient. The additional cathode flow allows more CO₂ to escape unless the cathode CO₂ concentration is simultaneously reduced.

Fig. 6 highlights methane composition in the biogas, which is known to fluctuate substantially. Simple fuel control methods using voltage and current measurements can ensure the anode chemical potential remains nearly constant without sampling the fuel composition. The concentration of diluents in the biogas, chiefly CO₂, can reduce system voltage while increasing CCR potential.

Fig. 5 illustrates how an increase in fuel CO₂ concentration (and thus a diminution of CH₄ concentration) increases CCR for the first configuration. This is understood since the biogas CO₂ remains entrained in the anode stream.

Fig. 7 illustrates the dependence of CCR on U₉CO₂ (as demonstrated in the Equation (2)). The CO₂ utilization factor in the first configuration is a function of the anode exhaust injection into the cathode stream. Any excess CO₂ injection escapes with the cathode exhaust. Fig. 6 also shows the linear relationship between CO₂ utilization and hydrogen recovery. The stack electrical efficiency

**Table 2**

Results (Configurations 1 and 2).

|                      | Configuration 1 | Configuration 2 |
|----------------------|-----------------|-----------------|
| Biogas composition (molar) | 0.65 CH₄–0.35 CO₂ | 0.65 CH₄–0.35 CO₂ |
| Flow rate Cathode/Biogas [kg s⁻¹] | 2.12/0.035 | 1.91/0.033 |
| Fuel Utilization, Uf (%)  | 65             | 65             |
| CO₂ utilization (%) – UCO₂ | 80             | 32.4           |
| Voltage [V]         | 0.677           | 0.733          |
| Current [mA cm⁻²]   | 177.5           | 163.1          |
| Plant Electric Efficiency — η | 0.322           | 0.362          |
| η with hydrogen (1HV) | 0.609           | 0.648          |
| H₂ produced [g s⁻¹] | 1.709           | 1.564          |
| Pₐ, stack [kW]      | 300             | 300            |
| Pₑ, plant [kW]      | 229.4           | 236.9          |
| CO₂ captured, vented [g s⁻¹] | 33.67–25.17  | 147.42–194.49 |
| Captured CO₂ purity (vol.) | 0.978         | 0.978          |
| CCR [%]              | 56.4            | 42.9           |
| Cathode inlet [CO₂] /[O₂] | 0.037/0.202  | 0.107/0.077   |
| Tₐ, inlet/Outlet [°C] | 550/650        | 550/650        |
| Tₑ, inlet/outlet [°C] | 530/612        | 530/622        |
| Pₑ, ICE [kW]        | —               | 1426           |

**Fig. 4.** U₀₂ — Voltage.

**Fig. 5.** Configuration 1, U₉ variation.

**Fig. 6.** Configuration 1, variation of methane composition in biogas.
slightly decreases as CO₂ utilization increases. Voltage and efficiency are affected by an increase in concentration overvoltage caused by limitations in CO₂ diffusion to the triple-phase-boundary. In particular, at high utilization the cathode CO₂ is close to stoichiometric and may be completely depleted in some cathode channels unless there is perfect manifolding and distribution of the oxidant stream. With stack power held constant at 300 kW, the current increases proportional to any voltage reduction. Oxygen utilization decreases with the rising U<sub>CO₂</sub> value, due to the additional air flow requirement for cooling. Lower electrochemical efficiency implies additional internal heat generation, and consequently rising air flow or rising temperature.

5.3. Configuration 2: internal combustion engine exhaust

The second configuration developed applies the exhaust of a separate ICE as the cathode oxidant feedstock. This integration removes the requirements for an air blower and for anode exhaust injection into the cathode stream. Without a means for controlling the cathode inlet concentration of CO₂ it is not possible to control the U<sub>CO₂</sub> as in the first configuration. Without anode exhaust injection the ICE must be of sufficient size to meet the O₂ and CO₂ requirement of the MCFC electrochemistry. This corresponds to an ICE: MCFC size ratio of 1.9:1 under nominal operating conditions. The ratio of ICE to MCFC electrical output will change as the O₂ and CO₂ requirements change with MCFC operating conditions. The ICE exhaust serves the additional purpose of providing the stack cooling flow. A 40% efficient ICE, operating on natural gas, with an exhaust flow matching the cathode flow requirement of an MCFC will contain 250% of the necessary CO₂, more for bio-gas and nearly 400% of the necessary O₂ for the electrochemical reactions. This corresponds to an ICE: MCFC size ratio of 4.66:1. Where the CO₂ exhaust of the ICE determines the smallest ICE size without anode exhaust injection, the cooling flow requirement determines the largest ICE size for which all of the ICE exhaust can be supplied to the MCFC. Between these two sizes some form of cathode recirculation or series connection of MCFC must be employed with intercooling to provide sufficient stack cooling. The ratio of MCFC to ICE size will influence the net cost, efficiency, CO₂ recovery and operational flexibility. This analysis will focus on the efficiency and CO₂ capture.

Cathode recirculation could partially decouple the inlet composition from flow rate, but without an additional source of CO₂ it will only serve to dilute both the inlet oxygen and CO₂ concentrations. Thus, a practical limit of cathode recirculation when using ICE exhaust is 20%. The ability to recirculate cathode exhaust is necessary if the ICE is to operate in a load following manner. All other system components are the same as for the previous reference configuration. A standard plant could readily be switched between self-CO₂ injection and this externally sourced CO₂ injection method, since the configuration of heat exchangers is very similar.

The ICE exhaust composition is calculated starting from the same biogas concentrations specified for the fuel cell. The electric power supplied by the biogas driven ICE is specified by \( P_{el,ICE} \) and Equation (3). Since we are dealing most likely with large diesel ICE an engine efficiency of 40% is specified with the methane molar flow described by Equation (4) and Equation (5).

\[
P_{el,ICE} = \text{LHV}_{CH_4} \times \dot{N}_{CH_4,biogas} \times \eta_{ICE} \tag{3}
\]

\[
\dot{N}_{CH_4,biogas} = \dot{N}_{CO_2,exhaust} \times y_{CH_4} \tag{4}
\]

\[
\dot{N}_{CO_2,exhaust} = \dot{N}_{exhaust} \times [CO_2] \tag{5}
\]
The ICE exhaust molar flow that feeds the cathode is calculated by the model in order to maintain the temperature difference of 100 K between inlet and outlet, while the carbon dioxide concentration depends upon the biogas concentration and upon the air index.

The efficiency and exhaust composition define the ICE system output, which can be successfully integrated with the MCFC CO2 recovery system. The CCR for this configuration (Equation (6)) is calculated as the ratio of the carbon recovered to the carbon that enters in the system in the form of biogas fuel to both the ICE and MCFC. Fig. 3 also illustrates this second configuration when internal recirculation of CO2 is replaced with ICE exhaust.

\[ CCR = \frac{N_{\text{CO}_2\text{rec}}}{N_{\text{BIOGAS,in}} + N_{\text{BIOGAS,ICE}}} \] (6)

Since none of the CO2 participating in the electrochemistry is subsequently injected into the cathode and exhausted, as for the configuration 1, simulations indicate a fourfold increase in the amount of CO2 recovered despite the lower CCR value of 42.95% (Table 2). At this largest size ratio, 1.4 MW ICE paired with a 300 kW MCFC, there is substantial excess CO2 available resulting in a lower UCO2 and a potential co-benefit of higher cell voltage due.

5.4. Sensitivity analysis – Configuration 2

As in the previous configuration, Fig. 8 shows that fuel utilization within the MCFC can be readily controlled. The impact on CCR exhibits similar trends, albeit with less variation. The equivalent electric power increases with \( U_e \) due to the reduced parasitic of the HSU. The HSU parasitic is reduced because there is less hydrogen available to recover. Electric performance remains strongly dependent upon \( U_e \). The results indicate a co-production flexibility, as it is possible to alter the ratio of hydrogen and electric production without dramatically impacting CCR.

The impact of varying biogas composition was also investigated for this configuration (Fig. 9). The primary difference is that the same biogas now feeds both the MCFC and the ICE. Hydrogen recovery and plant electrical efficiency do not change with the biogas variation, while CCR increases slightly. In particular, an increase in methane concentration lowers the ICE exhaust CO2 concentration and subsequently increases UCO2. Moreover, feeding the plant with pure methane \((y_{CH_4} = 1)\) involves the utilization of an engine with a nominal electric power that is increased compared to the standard case \((y_{CH_4} = 0.65)\).

5.5. Configuration 3: multiple stacks in series

A smaller ratio of ICE to MCFC sizes can result in much higher CO2 recovery, but requires some form of cathode recirculation and intercooling to achieve sufficient stack cooling. This configuration will consider the same 1.4 MW ICE paired with multiple 300 kW MCFC systems connected in series with cathode exhaust heat recovery placed between them as illustrated in the Fig. 10.

The internal combustion engine feeds the first MCFC cathode which subsequently supplies the second and third MCFC. Each MCFC requires a similar quantity of biogas. Thus, the total biogas flow to the fuel cell is tripled. The electric output and hydrogen

| Cross-flow configuration | \( P_e \) [kW] | CCRTot | CCRSTACK | UCO2 | \( \Delta T_{\text{cath}} \) [K] | H2STACK | H2 Tot [g/s] |
|-------------------------|---------------|--------|----------|------|-----------------|---------|----------|
| 1^ MCFC                 | 300           | 0.419  | 0.419    | 0.316| 100             | 0.454   | 1.586    |
| 2^ MCFC                 | 300           | 0.734  | 0.5248   | 0.414| 101             | 0.438   | 1.604    |
| 3^ MCFC-A               | 300           | 0.984  | 0.773    | 0.689| 112             | 0.414   | 1.737    |
| 3^ MCFC-B               | 150           | 0.868  | 0.755    | 0.664| 110             | 0.415   | 0.874    |
recovery are similarly tripled. This solution does not add to the HSU or carbon recovery balance of plant as the separate anode streams can be readily combined into a single effluent and handled as in configurations 1 and 2. The first two cathode outlet flow streams are cooled before entering downstream cathode compartments.

The two obstacles of this solution are: 1) With additional MCFC in series the UCO2 of the downstream systems increases considerably and these may experience the performance loss as described in the first configuration, and 2) Independent control of the temperature gradient is complicated by the coupling of the cathode flow rates. This problem could be limited by adding an additional air flow stream, but this solution would further dilute the CO2 concentration.

For these reasons, a practical limit to this series configuration was selected as 3 systems in series. Additional series and parallel configurations are potentially possible depending upon the electrical power requirements and biogas availability.

The CCR for this series connected configuration is calculated using Equation (7):

$$CCR = \frac{\sum N_{CO2\text{out}}}{N_{Biogas\text{to}\_1\text{MCFC}} + N_{Biogas\text{to}\_2\text{MCFC}} + N_{Biogas\text{to}\_3\text{MCFC}} + N_{ICE}}$$

Table 3 summarizes the primary performance characteristics of the series connected configuration. The first row, 1st MCFC, represents the original configuration while the 2nd and 3rd rows represent two and three systems connected in series. The High UCO2 in the third fuel cell resulted in significant performance loss, thus a final configuration of 2.5 systems in series was considered and presented in the last row of Table 3.

The two issues described previously, the progressive increase of UCO2 and cathode ΔT, are apparent in the increased heat generation of the downstream MCFC. The temperature gradient is constrained by materials constraints, and thus exceeding this limit can be dangerous for the life of the stack. The third MCFC would be expected to be the most likely to exceed temperature limits due to the increased UCO2 and fixed cathode flow. This can be solved by operating downstream MCFC at reduced power to maintain the same voltage as the upstream systems. The CCR in a series connected system exceeds 73% for two systems in series and exceeds 90% for 3 systems in series, when operated in this manner. This remarkably high carbon recovery is possible with the majority of the power still delivered by the low cost ICE system, 1.4 MW of 2.3 MW total. The system retains its hydrogen co-production capability and offers additional opportunity for high quality waste heat recovery between the high temperature fuel cells.

6. Summary and conclusions

Three possible configurations were outlined and studied; a standalone MCFC tri-generation plant, the same plant integrated with exhaust from an ICE plant, and the ICE-MCFC hybrid integrated with additional MCFC plants in a series configuration. The systems described could be readily deployed using existing commercial systems. Moreover, the first and the second configurations, while appearing more complex, could be realized in a single system with some valve design and controls for switching between ICE exhaust and CO2 injection from anode recirculation. A series configuration of MCFC technology coupled with an ICE achieves outstanding carbon recovery (exceeding 90%), with minimal parasitic load; that of the production of O2 in an ASU and the cooling fan in the second condenser.

Carbon separation and hydrogen co-production processes are compatible and benefit from the carbonate ion charge carrying properties of a molten carbonate fuel cell. Hydrogen co-production reduces fuel-to-electric efficiency due to the PSA parasitic load and lower operating fuel utilization used to generate excess hydrogen. All three configurations achieved notable carbon removal with minimal parasitic load. Configuration 1 (standalone system) required a high CO2 utilization factor (>75%) in order to separate a sufficient carbon quantity. Possible drawbacks to carbon recovery from a MCFC tri-generation system include complex thermal integration due to additional heat exchange steps as well as reduced fuel-to-electric efficiency due to the slow fuel utilization and high parasitic electric load of the HSU and any additional compression of CO2 and H2 products not considered in this analysis.

Appendix A: Cornerstones for the Configurations 1 and 2

| Table 4 | Cornerstones (Configuration 1). |
|---------|---------------------------------|
| T [˚C]  | XCH4  | XCO  | XCO2 | XH2 | XH2O | XH2i | XH2o | XCO2 | N [Mol s⁻¹] | m [kg s⁻¹] |
| 1       | 27    | 0.038 | 0.001 | 0.790 | 0.210 | 73.655 | 2.124 |
| 2       | 26    | 0.038 | 0.001 | 0.760 | 0.202 | 76.602 | 2.252 |
| 3       | 550   | 0.008 | 0.001 | 0.796 | 0.196 | 73.141 | 2.114 |
| 4       | 651   | 0.008 | 0.001 | 0.796 | 0.196 | 73.141 | 2.114 |
| 5       | 159   | 0.008 | 0.001 | 0.796 | 0.196 | 73.141 | 2.114 |
| 6       | 27    | 0.650 | 0.350 | 1.365 | 0.035 | 2.219 | 0.040 |
| 7       | 234   | 0.133 | 1     | 3.584 | 0.075 | 3.584 | 0.075 |
| 8       | 387   | 0.248 | 0.619 | 7.667 | 0.214 | 7.667 | 0.214 |
| 9       | 530   | 0.248 | 0.619 | 7.667 | 0.214 | 7.667 | 0.214 |
| 10      | 612   | 0.050 | 0.050 | 0.409 | 7.667 | 0.214 | 7.667 | 0.214 |
| 11      | 350   | 0.040 | 0.112 | 0.370 | 7.667 | 0.214 | 7.667 | 0.214 |
| 12      | 361   | 0.011 | 0.151 | 0.370 | 7.667 | 0.214 | 7.667 | 0.214 |
| 13      | 199   | 0.011 | 0.151 | 0.370 | 7.667 | 0.214 | 7.667 | 0.214 |
| 14      | 206   | 0.003 | 0.244 | 0.024 | 5.012 | 0.166 | 5.012 | 0.166 |
| 15      | 10    | 0.005 | 0.159 | 0.362 | 7.667 | 0.214 | 7.667 | 0.214 |
| 16      | 10    | 0.005 | 0.088 | 0.028 | 4.157 | 0.164 | 4.157 | 0.164 |
| 17      | 10    | 0.005 | 0.878 | 0.002 | 1.94 | 0.006 | 1.94 | 0.006 |
| 18      | 10    | 0.005 | 0.115 | 0.001 | 4.141 | 0.169 | 4.141 | 0.169 |
| 19      | 10    | 0.005 | 0.017 | 0.001 | 3.729 | 0.162 | 3.729 | 0.162 |
| 20      | 10    | 0.005 | 0.017 | 0.001 | 3.729 | 0.162 | 3.729 | 0.162 |
| 21      | 10    | 0.005 | 0.017 | 0.001 | 3.729 | 0.162 | 3.729 | 0.162 |
| 22      | 10    | 0.005 | 0.017 | 0.001 | 3.729 | 0.162 | 3.729 | 0.162 |
The Nernst potential is related to the Gibbs energy by the Faraday’s law and the number of electrons consumed in the reduction reaction (Equation (B4)). Nernst potential is calculated locally for the resistance proposed by Rivera[23].

\[ \eta_{conc} = \left( 1 + \frac{1}{\alpha} \right) \frac{RT}{nF} \ln \left( \frac{i_L - i}{i_L} \right) \]  

**Appendix B: overvoltage and reforming equations**

**Nernst potential and overvoltage losses**

The Nernst potential is related to the Gibbs energy by the Faraday’s law and the number of electrons consumed in the reduction reaction (Equation (B1)). Nernst potential is calculated locally using local bulk flow species concentrations and temperatures.

\[ E = \frac{\Delta G^0}{2F} - \frac{RT}{2F} \ln \left( \frac{p_{aH_2}p_{aCO_2}P_{aH_2O}^{0.5}}{p_{cH_2}p_{cCO_2}} \right) \]  

(B1)

In order to obtain the local voltage of the cell, it is necessary to subtract the three overvoltage losses (e.g. \( \eta_{act}, \eta_{ohm}, \eta_{conc} \)) from the Nernst potential. The expressions utilized to calculate the activation, ohmic, and concentration losses are detailed in Equations (B2)–(B4). An equipotential constraint is used to solve for the spatial current distribution which balances the temperatures and concentration variation. The activation overvoltage, \( \eta_{act} \), is related to the charge transfer from the molecule to the electrode (oxidation) and vice versa (reduction). Equation (B2) relates the activation polarization and the current density through the Tafel’s law, where \( \eta = 0 \) is the exchange current density (300 A m\(^{-2}\)) and \( \alpha \) is the charge transfer coefficient (0.4).

\[ \eta_{act} = \frac{RT}{nF} \ln \left( \frac{i}{i_0} \right) \]  

(B2)

The ohmic overvoltage, \( \eta_{ohm} \), is significant due to the poor electrical conduction of the electrodes. The conduction is highly temperature dependent. This model utilized an empirical expression for the resistance proposed by Rivera [23].

\[ \eta_{ohm} = i \cdot \left( 4.7833 \times 10^{-4} - \left( 6.6667 \times 10^{-7} \right) (T - 273) \right) \]  

(B3)

The concentration overvoltage, \( \eta_{conc} \), is associated with diffusion of bulk reactants to the triple phase boundary. The diffusion losses are modeled with a logarithmic function with an empirically identified current density limit, \( i_L \) (6000 A m\(^{-2}\)).

\[ \eta_{conc} = \left( 1 + \frac{i}{i_L} \right) \frac{RT}{nF} \ln \left( \frac{i_L - i}{i_L} \right) \]  

(B4)

\[ \frac{dX_{i, out}}{dt} = \frac{R_{REFORM} + R_{WGS} + R_{CONSUME} + N_{in}X_{i,in} - N_{out}X_{i,out}}{P_{i, f}} \]  

(B5)

The rate of species conversion within the reformer and anode channels, \( R_{REFORM} + R_{WGS} \), is determined from the chemical kinetics of steam methane reforming and water gas shift reactions. Other chemical reactions can take place, but they occur at a much slower rate which allows a model to maintain accuracy when neglecting them. The methane reforming rate uses an empirical correlation (Equation (B6)) developed by Seo et al. [24]. The water gas shift reaction (Equation (B7)) has been adopted from Haberman and Young [23], and was developed specifically for the high temperature conditions of a fuel cell anode.

\[ R_{REFORM} = A_{cell} \cdot p_{CH_4} \cdot \exp \left[ - \frac{8.2 \times 10^4}{R_0 T} - 2137 \times 10^6 \right] \]  

(B6)
\[ R_{\text{WGS}} = A_{\text{cell}}' P_{\text{CO}} \times 0.0171 \exp \left( \frac{-103191}{R_a T} \right) \times \left( 1 - \frac{P_{\text{CO}} P_{\text{H}_2}}{\exp \left( \frac{4276}{T} - 3.196 \right) P_{\text{CO}} P_{\text{H}_2}} \right) \]  \tag{B7}

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