CO₂ Utilization Technologies: A Techno-Economic Analysis for Synthetic Natural Gas Production

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Abstract: Production of synthetic natural gas (SNG) offers an alternative way to valorize captured CO₂ from energy intensive industrial processes or from a dedicated CO₂ grid. This paper presents an energy-efficient way for synthetic natural gas production using captured CO₂ and renewable hydrogen. Considering several renewable hydrogen production sources, a techno-economic analysis was performed to find a promising path toward its practical application. In the paper, the five possible renewable hydrogen sources (photo fermentation, dark fermentation, biomass gasification, bio photolysis, and PV electrolysis) were compared to the two reference cases (steam methane reforming and water electrolysis) from an economic stand point using key performance indicators. Possible hydrogen production capacities were also considered for the evaluation. From a technical point of view, the SNG process is an efficient process from both energy efficiency (about 57%) and CO₂ conversion rate (99%). From the evaluated options, the photo-fermentation proved to be the most attractive with a levelized cost of synthetic natural gas of 18.62 €/GJ. Considering the production capacities, this option loses its advantageousness and biomass gasification becomes more attractive with a little higher levelized cost at 20.96 €/GJ. Both results present the option when no CO₂ credit is considered. As presented, the CO₂ credits significantly improve the key performance indicators, however, the SNG levelized cost is still higher than natural gas prices.

Keywords: CO₂ utilization; synthetic natural gas; techno-economic analysis; renewable hydrogen

1. Introduction

The latest Intergovernmental Panel on Climate Change (IPCC) special report on global warming of 1.5 °C [1] urges for the reduction of fossil CO₂ emissions in order to limit global warming and to reduce the risks associated with a higher temperature increase caused by the growing atmospheric CO₂ concentration. Global population and economic growth go hand-in-hand with higher energy intensity, which also mean higher anthropogenic fossil CO₂ emissions. In the past years, a decoupling trend has been observed between the economic growth and greenhouse gas emissions [2], however, to reach the goals set within the 1.5 °C model, net zero emissions have to be reached by 2050 [1]. The International Energy Agency (IEA) reemphasized the importance of carbon capture, utilization, and storage (CCUS) in the process of lowering anthropogenic CO₂ emissions [2], however, this will result in a significant abundance of CO₂ in the near future. Available geological CO₂ storage sites are distributed unevenly geographically. In this case, two options are possible: either set up a CO₂ transport system capable of transporting high quantities of CO₂ from one place to another as presented by Reiter et al. [3], or build CO₂ utilization facilities that can chemically convert the captured CO₂ into value added products (e.g., chemicals, energy carriers). The production of synthetic natural (SNG) is a noteworthy option of CO₂ utilization that has a high hydrogen demand. The hydrogen demand required for SNG production is addressed in the next section of the paper.
1.1. Hydrogen Production Options

Steam methane reforming (SMR) is currently the most important technology for hydrogen production [4] using fossil fuels (natural gas or light hydrocarbons). Considering the high abundance of these fossil sources and the overwhelming dominance of this technology in hydrogen production, this can be expected to stay the same in the short to medium future. In light of the CO$_2$ mitigation goals, renewable-based hydrogen production coupled with CO$_2$ capture and utilization can be a viable option for the transition into a low carbon hydrogen-based society [5]. Less carbon intensive hydrogen production options are available in the literature, however, as the production of renewable hydrogen is energy intensive, the production costs are also high. The use of renewable power for hydrogen production is one of the most important alternatives, and depending on the carbon intensity of the power produced, one can assume a similar degree of CO$_2$ emissions when comparing fossil fuel reforming technologies and power-based technologies.

On the other hand, several carbon free hydrogen sources exist that can produce hydrogen with low-to-near-zero emissions. These hydrogen production options tend to be less energy efficient technologies, however, these technologies need to be further improved. In the following section, aside from the five evaluated renewable hydrogen production options, the two reference hydrogen production technologies (steam methane reforming and water electrolysis) are presented in brief.

Steam methane reforming is one of the reference hydrogen production technologies considered in this paper. This involves the reacting of a hydro-carbonated fossil fuel, generally natural gas (NG) with steam in the presence of a Ni-based catalyst, to yield hydrogen according to Equation (1) (together with water gas shift—see Equation (4)).

$$CH_4 + H_2O \leftrightarrow 3H_2 + CO$$

The reforming reaction is highly endothermic ($\Delta H_r = + 206 \text{ kJ/mol}$) and it is carried out commonly in an externally heated reactor at high temperature ($500–900$ °C) and at a pressure range of 20–35 bar. Other technological options are autothermal reforming or partial oxidation. Steam-to-carbon ratio is an important parameter of the process, usually this has value of 2.5:3. Low pressure, high temperature, and high steam-to-carbon ratio favors the formation of hydrogen. The high steam-to-carbon ratio also limits the deposition of carbon of the catalyst; the deposition reaction being a major problem in reforming processes. A Ni-based catalyst is applied for the reforming process [4] thanks to its high activity and low cost, however, this needs special attention because of its sensibility to sulfur (poisoning). SMR is the most widely used reforming process for hydrogen production, and autothermal reforming and partial oxidation are two major variants of the reforming process, the differences being that the first is an autothermal process while the second is an exothermic process, respectively [6].

Biomass gasification is an important process that gives syngas, a mixture of CO and H$_2$. With this technology, through specific separation techniques (using gas-liquid and gas-solid systems), high purity hydrogen can be obtained from any solid fuel (either fossil or renewable). For the purpose of this paper, the gasification of biomass was considered. In the process, oxygen and steam were reacted with the biomass (e.g., sawdust, agricultural wastes, etc.) through a series of reactions that deliver hydrogen, carbon monoxide, and methane aside from other minor components as presented in Equations (2) and (3).

$$\alpha C_k H_l O_m + \beta H_2O \xrightarrow{\text{heat}} bCO + cCO_2 + dCH_4 + eH_2 + fTar$$

$$C_x H_y + xH_2O \xrightarrow{\text{heat}} \left(\frac{y}{2} + x\right)H_2 + xCO$$

In Equation (2), $C_k H_l O_m$ represents the general formula of the biomass that can be used for the production of hydrogen. This could result from wood, sawdust, sugar cane, or any other agricultural waste. Tar is an undesired product of the gasification reaction that causes slugging and fouling. To counteract its formation, a good catalyst is required
that stops the formation of side reactions. During gasification, hydrogen is obtained by the general Equations (2) and (3) as well as from the water-gas shift (WGS) reaction (presented in Equation (4)). The WGS reaction is used to convert the produced CO to CO₂ while additional hydrogen can be obtained as presented in the equation. In order to convert as much CO as possible, an additional unit is added to the system.

\[ CO + H_2O \leftrightarrow H_2 + CO_2 \]  

The most well-known gasification processes are licensed by General Electric (Massachusetts, USA), Siemens (Berlin, Germania) and Shell (London, England). All of these are entrained-flow gasifiers, meaning that the solid fuel is fed into the reactor using an inert transport gas (N₂, CO₂) or water slurry. The temperatures inside the gasifier can reach up to 1350–1500 °C whereas the pressure can go up to 100 bar [7]. The operating parameters of the gasifier not only greatly depend on the fuel and the produced ash, but also on up-stream and down-stream units.

**Electrolysis** and photovoltaic (PV) electrolysis uses water and power for the production of hydrogen. It is regarded as the most basic industrial technology for high purity hydrogen production. An electrical current is passed through liquid water between two electrodes while oxygen and hydrogen gasses are obtained. The two gasses are easily separated as they are formed on the two electrodes: hydrogen on the cathode and oxygen on the anode. This is the major advantage of the process: it offers a low-carbon (provided that carbon-free electricity is used) hydrogen production technology with the only side product being oxygen. Using power from regenerable sources (PV energy for the case of PV electrolysis), this can reduce the CO₂ impact of the process to near zero. The major drawback of water electrolysis is its low energy efficiency, which explains the low share of electrolysis in global hydrogen production of about 4% [5]. This low efficiency also translates into high economic costs with the operation of electrolysers [8]. On the other hand, this share of hydrogen produced via electrolysis is expected to increase in the future as the need for clean hydrogen will increase, but also as a new class of electrolysers are developed (solid oxide electrolysers). Table 1 presents a comparison between the three most important electrolysis technologies: alkaline water electrolysis, polymer membrane electrolysis, and solid oxide electrolysis. Although rather new, solid oxide electrolysis shows great promise in reducing the high costs associated with the process [9].

**Table 1.** Most important water electrolysis technologies.

| Technology Maturity | Alkaline | Polymer Membrane | Solid Oxide Electrolysis |
|---------------------|----------|------------------|-------------------------|
| Cell temperature (°C) | 60–80 | 50–80 | 900–1000 |
| Cell pressure (bar) | <30 | <30 | <30 |
| Current density (A/cm²) | 0.2–0.4 | 0.6–2.0 | 0.3–1.0 |
| Voltage efficiency (%) | 62–82 | 67–82 | 81–86 |
| Specific system energy consumption (kWh/Nm³) | 4.5–7.0 | 4.5–7.5 | 2.5–3.5 |
| Stack lifetime (h) | <90,000 | <20,000 | <40,000 |
| Cold start-up time (min) | 15 | <15 | >60 |

Electrolyzing units, in general, are highly sensitive to the purity of the water, for example, the presence of chlorine in the water will result in chlorine gas formation instead of oxygen. This problem can be solved by using water purifiers or special alloys, in the case of high salt content water as presented in the literature [10,11].

PV electrolysis uses renewable electricity produced by PV panels for the conversion of solar power into electrical power and from there to chemical energy by producing hydrogen. Although the system shows great promise, the overall energy efficiency is rather low when compared to other hydrogen production options. The PV system itself needs
several electrical components aside from the electrolysis cell such as the PV panels, DC bus bar, a link to the AC grid, and a set of batteries. For storage, a small sized hydrogen storage tank is also required. At this moment, this option has one of the highest costs, with the price of hydrogen produced being up to 25 times higher than fossil fuel alternatives (e.g., hydrocarbon catalytic reforming), however, this gap has been continuously dropping in the past years and this factor is expected to drop to six in the near future [12].

**Dark fermentation** is a biochemical process performed in the absence of light by microorganisms. The biochemical energy stored in organic matter is used by microorganisms to extract hydrogen from the biomaterial fed into the bioreactor. If light is excluded from the system, dark fermentation is favored, and by specific microorganisms, the organic matter is transformed, resulting in high energy gasses, with hydrogen gas being one of these possible products. The process therefore is an attractive green process for hydrogen production as it uses regenerative sources and offers a controllable microbial growth in waste processing. The process, aside from producing hydrogen, has the advantage that it offers an alternative to waste processing.

These microorganisms are categorized based on their temperature preferences. The processes that prefer medium temperatures are called mesophilic processes while microorganisms that prefer high temperatures are called thermophilic. Koutrouli et al. [13] evaluated the hydrogen production from olive pulp and showed that thermophilic hydrogen production was more efficient than mesophilic in both the hydrogen production rate and can yield a hydrogen production rate of 0.32 mole hydrogen per kg olive pulp solution. On the other hand, a thermophilic process requires higher costs for operation when compared to a mesophilic process. Similarly, high hydrogen production rates can be obtained in waste water treatment plants using anaerobic sequencing, as presented in the work of Thammanoon et al. [14]. The hydrogen production rate greatly depends on the system parameters and also from the substrate source, as highlighted by Giordano et al. [15].

**Photo-fermentation and bio-photolysis** processes are biochemical processes that use solar energy to decompose and synthesize organic material, respectively, while producing hydrogen. In contrast with dark fermentation, photo fermentation and bio photolysis use the energy of photons to also produce, aside from other compounds, hydrogen. The major difference between photo fermentation and bio photolysis is the starting material for the process. While bio photolysis uses inorganic materials as water, CO₂, and nutrients for the production of organic material, photo fermentation decomposes organic matter. Both processes produce hydrogen as a side product that can be separated and used as a fuel. Kotay et al. [16] grouped bio photolysis into two categories: direct photolysis and indirect photolysis, depending on whether the photons interact directly or indirectly (through a promotor molecule [17]) with the substrate. Both processes require only the microorganisms and sufficient sunlight, making it a viable option for renewable hydrogen production. However, two major drawbacks of the processes need to be addressed. The first is a high capital cost because of the high surface area photobioreactor and the low hydrogen production rates compared to competing technologies. Both drawbacks need to be reduced to maintain the economic attractiveness of these processes [16].

### 1.2. Power-to-Gas (PtG) Route

Synthetic natural gas can be obtained with the exothermic reaction of hydrogen with carbon dioxide (CO₂ methanation), as presented in Equation (5):

\[
\text{CO}_2 + 4\text{H}_2 \leftrightarrow \text{CH}_4 + 2\text{H}_2\text{O} \Delta H^0 = -165 \text{ kJ/mol} \quad (5)
\]

The process shows significant interest in light of the high share of variable renewable energy sources available at this moment with the aim of storing the surplus renewable energy using electrolyzers for producing hydrogen, which in turn, can be stored for longer periods and used as fuel when production is low. Storing is possible in several forms in the production of hydrogen [3] as this product has a high demand in the chemical market and is also expected to play an important role in the struggle to lower CO₂ emissions. Its major
drawback, however, is its low energy density when compared to other possible products such as methane and methanol. Methanol production from CO$_2$ is a good alternative, however, the potential market for it is significantly smaller than for methane.

The high methane demand drove this analysis to focus mainly on the production of synthetic natural gas, which lately has been of high interest [18,19]. This process can also be considered as power-to-gas technology, as the surplus power is stored in the form of chemical energy. The increased variable renewable energy output translates into a major interest in energy production optimization [20] and energy storage options [21], starting from surplus power in the national power grid. Gutiérrez-Martín et al. [22] presented a comprehensive study of the power-to-gas route including detailed economic calculations. In this work, usage of the surplus power available in the national power grid to power an electrolyzer for hydrogen production was considered. Considering several products as energy carriers (e.g., hydrogen, methane), the possibility of returning the power extracted and stored in the form of a gas to the power grid when power demand is high was evaluated. The paper shows promising results for the economics of the proposed plants, however, a major contributor to the cost of the energy obtained and stored in this manner greatly depends on the initial cost of the power used in the electrolyzer. In their calculations, they assumed free power from the peak-shaving. When a low electricity price is considered, production prices increase significantly. Buchholz et al. [23] used the power-to-gas route for storing surplus power in the form of synthetic natural gas combined with a lignite power plant to increase the flexibility in light of the variable output of renewables. The results show promising results from an economic perspective, however, as the power-to-gas route seems to resolve the issue with peak-shaving, as Vandewalle et al. [24] points out, this only transfers the strain from the power grid to the gas grid. Although at first glance this does not resolve the issue, the surplus of energy being more spread out in several sectors on a global scale acts as an equalizer of energy between sectors and lowers the effect of overproduction. Second, with a proper storage capacity, natural gas overproduction can be minimalized. Aside from the economic issues, another problem with SNG is the difference in composition and heating value from natural gas. This issue is discussed briefly in the next section of the paper.

2. Synthetic Natural Gas Production Process

2.1. Synthetic Natural Gas (SNG) Quality Requirements

Natural gas quality is an important factor when discussing synthetic natural gas production, however, this aspect tends to be overlooked in papers published on power-to-gas technologies [3,18,25]. In general, gas composition is used for the evaluation of natural gas proprieties. The major components of natural gas are hydrocarbon species (methane, ethane, propane etc.), the inert constituents are nitrogen and carbon dioxide, and undesirable species such as the sulfur containing components, water, and mercury. For some natural gas users in the chemical industry, a specific composition has to be tuned, however, for the majority of gas users, combustion characterizing parameters are sufficient like the Wobbe Index (WI), the calorific value, and the Methane Number (MN).

These parameters are monitored to ensure the safe and energy efficient operation of the plants. For gas transportation between countries and between quality zones, natural gas specifications are used from Directive 2003/55/CE [26]. As presented in the directive, the obtained NG methane content has to be a minimum of 95 vol.% and 90 vol.% for the high calorific gas grid and the low calorific gas grid, respectively, whereas the hydrogen content has to be limited to 2 vol.% if a natural gas filling station is linked to the grid as stated in the DIN EN 51624 standard [26].

The Wobbe Index (WI) is an important gas quality indicator in the case of most gas-burning equipment. A constant WI of the fuel results in a constant energy supply to the installation for a given temperature and pressure. The same WI can be obtained if the composition of the gas varies. Changes in the WI have an effect on the power output of the system and also on the air-to-fuel ratio. Most natural gas fueled units can compensate for
WI variations using feedback control, but only to a certain extent and always afterward. Fast variation in WI could lead to high ramps in power output, therefore, these need to be avoided because of the nature of the feedback control system [27].

Another important aspect as mentioned before is the hydrogen content of the NG. Hydrogen addition to the natural gas grid reduces the energy transportation of the pipeline as hydrogen has a lower compressibility as natural gas [28]. This is a non-linear effect that greatly depends on the total pressure of the gas and the partial pressure of hydrogen. One way to counteract this effect is by increasing the gas flowrate. Even so, limiting the hydrogen content to a certain value is highly recommended.

Finally, the operating pressure of the gas grid also has to be considered in the design phase of the plant. Each gas grid has a designated operating pressure interval that mostly depends on the number of consumers that are linked to it, the material of the pipes, and the estimated flowrate of gas. This way, within one grid, several pressure intervals are used. For instance, for the Romanian natural gas grid for household use, the operating pressures of the gas grid are presented in Table 2.

| Code Name | Pressure Interval (bar) |
|-----------|-------------------------|
| PJ        | ≤1.05                   |
| PR        | 1.05 < PR ≤ 3           |
| PM        | 3 < PM ≤ 7              |
| PI        | 7 < PI ≤ 11             |
|           | with special regulations also possible | >11 |

In the selection of the output pressure of the SNG system, one has to consider the designated market and the production rate. These pressures can range from the values presented in Table 2 up to 70 bar [30].

2.2. Process Limitations of CO₂ Utilization Technologies

The renewable hydrogen production options were compared to two carbon intensive options (steam reforming and water electrolysis). Therefore, the selection and design of the hydrogen production unit requires care. Hydrogen is assumed to be generated in place or in close proximity of the plant and this unit would serve the sole purpose of generating the gas required for the plant using the selected option. With this in mind, when selecting the geographical location of the plant, one has to consider the source of the starting material for the hydrogen production unit (biomass or solar energy) as well as the disposal of the side products generated. The transportation of these materials was not considered in this work. The feed parameters were assumed as the general parameters applied in hydrogen production plants as presented in Table 3.

| CO₂      | H₂      |
|----------|---------|
| Temperature | 15 °C   | 15 °C   |
| Pressure  | 100 bar | 60 bar  |
| Purity    | 99 vol.%| 99 vol.%|

Several studies have been published on assessing the CO₂ sources of a country and on the construction of a special grid for captured CO₂. Karjunen et al. [31] presented a case study for Finland focusing on the basic elements of the grid. Reiter et al. [3] presented a case study for Austria that focused on using the CO₂ generated in several parts of the country for a power-to-gas model through a CO₂ grid. As pointed out in the work, CO₂ is generated in high quantities at one point, and this might need to be transported for its use somewhere else. Therefore, CO₂ transportation is a complex phenomenon and a lot of work has to be put into the subject to be able to design a CO₂ grid capable of accepting
the CO₂ generated in one place and transport it to consumer facilities. In this work, it was assumed that the CO₂ is transported from a great distance and the feed-in parameters are selected according to the calculations of Karjunen [31], as presented in Table 3.

3. Techno-Economic Assessment Methodology

Equation (5), also known as the Sabatier reaction, is the catalytic methanation of CO₂, which is an equilibrium reaction depending on the conditions of the reaction. High pressure and low temperature favor the formation of methane. Figure 1 shows the influence of operating temperature (a) and pressure (b) on the gas composition resulting from the CO₂ methanation process.

![Figure 1. Influence of temperature (a) and pressure (b) to CO₂ methanation process.](image)

In the reaction system, the water gas shift (WGS) and CO methanation reactions also have a great influence on the product composition. The WGS and the CO methanation reactions are presented in Equations (6) and (7), respectively.

\[
\text{CO} + \text{H}_2\text{O} \leftrightarrow \text{H}_2 + \text{CO}_2 \Delta H^0 = -41 \text{ kJ/mol} \\
\text{CO} + 3\text{H}_2 \leftrightarrow \text{CH}_4 + \text{H}_2\text{O} \Delta H^0 = -206 \text{ kJ/mol}
\]

Catalysts applied for the process vary from noble metals to common nickel [32] and they can be applied in the temperature range between 250 °C and 550 °C [33]. Several pilot units for CO₂ hydrogenation were built in the past years as reported by Koytsoumpa and Karellas [26]. In the present work for the reaction conditions, an operating pressure of 50 bar and temperature of 300 °C were selected using a Ni-based catalyst according to the results of Chiang et al. [32].

3.1. Key Design Assumptions, Aspen Plus Modeling, and Thermal Integration Analysis

The conceptual design of a SNG plant is presented in Figure 2. In the case of an economic analysis, the estimation of capital and operational costs as well as SNG production cost are major components that depend on several factors (technological readiness, geographical location, etc.). As a system boundary, hydrogen production from various renewable sources, SNG plant, and its additional units were considered in the present analysis. We did not consider the CO₂ capture process either from energy efficiency point of view or from an economic cost implications point of view. In the present paper, to obtain an accurate estimation of the techno-economic parameters, process modeling and simulation using Aspen Plus® (Bedford, MA, USA) [34] software was used to generate the overall mass and energy balances. The key design parameters are presented in Table 4.
Figure 2. Proposed configuration for the synthetic natural gas (SNG) plant.

Table 4. Key modeling parameters applied in Aspen Plus®.

| Parameter                              | Value                  |
|----------------------------------------|------------------------|
| Phase equilibrium model                | Non-random two-liquid model (NRTL) |
| Methanation reactor model              | Gibbs reactor          |
| Operational parameters of methanation reactor | 300 °C, 50 bar       |
| Live steam properties                  | 280 °C, 3 bar          |
| Steam turbine efficiency               | 87%                    |
| Steam turbine condensation pressure    | 50 mbar                |
| Pump efficiency                        | 85%                    |
| Heat exchanger min. temperature difference | 10 °C             |
| Inlet/outlet cooling water temperature | 15 °C/25 °C            |

Hydrogen and carbon dioxide were fed into the system at the pressures corresponding to the transport parameters presented earlier, however, as these values were higher than the reactor pressure (especially, in the case of the CO₂) this had to be reduced. In order to reduce the ancillary power consumption and the operational costs, a turbine was placed to recover part of the energy released by decompression. The methanation reactor operated at 50 bar, as presented in Table 4. High pressure was selected to push the reaction toward the formation of methane according to Le Chatelier’s principle (see also Figure 1). The CO₂ methanation reaction was modeled using the thermodynamic data of the process simulator and compared to the literature data provided by Koytsoumpa et al. [26]. The CO₂ methanation reaction is highly exothermic, and the generated heat is removed by the generation of steam within the reactor. Through water evaporation, a constant temperature can be maintained in the reaction media. The generated steam is used in a steam turbine for power production (to cover the plant ancillary consumption as well as export to the grid).

The resulting gas stream from the CO₂ methanation process is cooled to 25 °C in several heat exchanger units to recover the available heat and to increase the overall plant energy efficiency. The condensed water is separated and can be recycled to the electrolyzer in the case that an electrolyzer is used for hydrogen production. The modeling of the electrolyzer was not included in this work. The outlet gas can be decompressed to meet the required pressure of the gas grid or can be stored in a special SNG tank for power production during high power prices. The water produced in the CO₂ methanation process leaves at a pressure of 50 bar. Additional energy can be recovered from this residual water stream by its decompression, however, this was not included in the model.

Pinch analysis is a powerful tool to evaluate the overall system energy integration, increase the overall energy efficiency, and minimize energy consumption by developing improved design alternatives. This tool was applied for the newly proposed SNG production.
plant. The hot and cold composite curves are presented in Figure 3. The heat removed from the CO₂ methanation reactor (about 65 MWth) was used for steam generation and then power production. The steam generation can be observed in Figure 3 by the horizontal segment of the cold composite curve (in blue).

![Figure 3. Hot and cold composite curves obtained on the proposed plant.](image)

The outlet gas stream leaving the CO₂ methanation reactor is cooled in a series of heat exchangers while steam is generated and then superheated. As presented by the hot composite curve (HCC), the product gas is cooled from 300 °C to 240 °C when the condensation of water starts at 50 bar. As the temperature drops to 180 °C, a significant part of the water has condensed. The remaining available heat is extracted by cooling of the gasses and the liquid water. The cold composite curve (CCC) represents the behavior of the water (and then steam) in the steam cycle. Heating starts from 33 °C to 140 °C where evaporation starts. After all the water has evaporated, the saturated steam is superheated to 280 °C.

As key technical performance indicators to be calculated from the mass and energy balances derived from simulation, the following elements were used: (i) total electricity input for hydrogen production (by water electrolysis considered as an illustrative example) was calculated as hydrogen mass flow multiplied by the specific electricity consumption of the electrolyzer; (ii) SNG thermal output was calculated as SNG mass flow multiplied by the correspondent SNG lower calorific value; (iii) total power production of the plant was calculated as electricity produced by steam cycle and gas decompression processes; (iv) ancillary power consumption of the plant was calculated by adding together all electricity consumptions of the plant sub-systems; (v) net energy efficiency represents the ratio between thermal energy of the SNG to the overall energy (electricity) consumption of the plant; and (vi) CO₂ conversion ratio shows the ratio of input carbon found in SNG.

### 3.2. Economic Assessment Methodology and Main Assumptions

Economic performance of the CO₂ hydrogenation process mostly depends on the cost of the hydrogen used as reactant. The same applies for this work, hence the importance of the comparison between several renewable hydrogen production sources. The costs of production for the evaluated hydrogen production technologies were obtained from the work of Dincer et al. [35] (see Table 5). Compared to these assumed hydrogen production costs, other relevant studies [5–7] showed relatively minor variations ± 10%. In any case, in the Results and Discussion section, a sensitivity analysis was done including hydrogen price variations.
Steam methane reforming was added to the model to have a base case, as hydrogen from NG is the leading hydrogen production method globally today. An economic model was also generated in the case of the standard electrolysis process, serving as a base case model for the PV electrolysis model. Photo fermentation, dark fermentation, biomass gasification, bio photolysis, and PV electrolysis were the evaluated renewable hydrogen sources in this paper. These technologies have roughly the same hydrogen cost (except PV electrolysis), which can be explained by the fact that the same low-cost starting material can be used. The production costs of renewable sources for hydrogen is about three to four times higher, meaning in order for the technologies to stay competitive, additional revenue sources need to be found. To understand the behavior of the PV electrolysis system, a standard electrolysis was added. By default, water electrolysis is a high cost technology, however, when the power needed is from PV panels, the cost of hydrogen doubles, making it the most expensive hydrogen source considered in this paper (4.75 €/kg).

### 3.2.1. Capital Cost Estimation

The capital cost of the units in the SNG plant were estimated using the methodology presented by Turton et al. [36]. The calculation methodology used for the estimation of the capital cost is presented in this section. In the first part of the calculation, the equipment purchase cost is calculated using Equation (8):

$$\log_{10} C_p = K_1 + K_2 \times \log_{10}(A) + K_3 \times \left[\log_{10}(A)\right]^2$$

(8)

In Equation (8), $C_p$ is the purchase cost; $K_1$, $K_2$, and $K_3$ are module specific constants; and A is a module specific parameter. Using the purchase cost, the bare module cost is obtained using Equation (9):

$$C_{BM} = C_p \times (B_1 + B_2 \times F_M \times F_P) = C_p \times F_{BM}$$

(9)

In Equation (9), $C_{BM}$ is the bare module cost; $B_1$ and $B_2$ are unit specific constants (Turton et al. [36]); $F_M$ is the material factor; and $F_P$ is the pressure factor. The constants applied for the capital cost estimation in Equations (8) and (9) are presented in Table 6. In the case of the heat exchangers and pressure vessels, the first part of Equation (9) is used whereas in the case of the decompressors and the pumps, the second part is used.

### Table 6. Module specific parameters and the constants applied for the capital cost estimation according to Turton et al. [36].

| Process Unit   | Module Specific Parameter | $K_1$  | $K_2$  | $K_3$  | $B_1$ | $B_2$ |
|---------------|---------------------------|-------|-------|-------|-------|-------|
| Heat exchanger | Heat exchanger area       | 4.3247| −0.303| 0.1634| 2.25  | 1.82  |
| Decompressor  | Fluid power               | 2.7051| 1.4398| −0.1776|       |       |
| Pump          | Shaft power               | 3.3892| 0.0536| 0.1538|       |       |
| Pressure vessel| Volume                    | 3.4974| 0.4485| 0.1074| 2.25  | 2.25  |

In the case of the heat exchangers and pressure vessels, the first part of Equation (9) is used whereas in the case of the decompressors and the pumps, the second part is used.
The USD to EURO conversion factor applied in the calculation was 1.2 $/€. The install
cost was obtained using an install factor of 1.68 and an up-to-date cost was obtained for
2020 using the Chemical Engineering Plant Cost Index (CEPCI) [37] using Equation (10):

\[
\frac{C_1}{C_2} = \frac{\text{Index}_1}{\text{Index}_2}
\]  

(10)

For the calculation of the total overnight cost, the European Benchmark Task Force
(EBTF) guidelines [38] were applied as this is a reliable option in the case of European
thermal power plant cost estimations. Table 7 presents the methodology applied for the
estimation of the total overnight cost (TOC).

| Component | Definition |
|-----------|------------|
| Bare erected cost (BEC) | Install cost of each unit |
| Engineering procurement and construction costs (EPCC) | 10% of BEC |
| Project contingency (PC) | 15% of (BEC + EPCC) |
| Total plant costs (TPC) | BEC + EPCC + PC |
| Owners cost (12% of TPC) | 12% of TPC |
| Total overnight costs (TOC) | TPC + Owners costs |

Engineering procurement and construction cost was estimated as 10%, a lower value
as the plant itself is a simple construction with no complicated processes, however, as
the plant can be considered as an emerging technology a project contingency of 15% was
applied. The rest of the costs were selected according to the general recommendations.

3.2.2. Operations and Maintenance (O&M) Costs

Table 8 presents the main assumptions used for calculation of fixed and variable
operations and maintenance (O&M) costs.

| Fixed Operations and Maintenance (O&M) Costs | Value | U.M. |
|--------------------------------------------|-------|------|
| Operating labor | 18.01 [39] | €/h |
| Maintenance, support and administrative labor | 2.0 | % of TOC |
| Cost of electricity | 85 [39] | €/MWh |
| **Variable O&M Costs** | | |
| Process water costs | 3.1 [39] | €/t |
| Cooling water make up costs | 2.5 [39] | €/t |
| Catalyst cost | 12,500 [40] | €/t |

A small number of SNG plant operators were selected, thanks to the simple plant
design; a total number of six persons per shift was considered, working in four shifts.
The total cost associated with the plant personnel was evaluated based on the hourly
cost of the work force as presented in Table 8. Similarly, literature data were used for the
rest of the operating and maintenance costs. For the surplus of power produced by the
plant, a high selling price was assumed as this can be considered as a renewable power
source to account for the credits resulting from producing renewable power. In the case of
the catalyst used in the CO₂ methanation reactor, a replacement period of two years was
assumed in accordance to the manufacturer’s specifications.

3.2.3. Cash Flow Analysis

To determine the key performance indicators (KPI) and obtain the levelized cost
of SNG, a discounted cash flow analysis was performed according to the description of
Smith et al. [41] on the process and compared to the present NG prices. In the analysis, all
the costs presented in the Methodology section were considered with an average capacity factor of 90%. The levelized cost (LC) of SNG was calculated considering a net present value (NPV) of zero at the end of the economic lifetime. The formula used for the calculation is presented in Equation (11).

\[
NPV = \sum_{t=0}^{n} \frac{ACF_t}{(1 + i)^t}
\]

where \( ACF \) is the annual cash flow; \( "i" \) is the interest rate; \( "t" \) is the actual year; and \( "n" \) is the economic lifetime. Table 9 presents the main assumptions for the cash flow analysis performed on the SNG production plant.

Table 9. Assumptions for cash flow analysis.

| Economic lifetime | 25 years |
|-------------------|----------|
| Interest rate     | 8%       |
| Construction period| 2 years |
| Capacity factor   | 90%      |
| First year capacity factor | 65%    |

The obtained levelized cost for the SNG was compared to the values presented in the relevant literature sources (e.g., IEAGHG report [4] on steam reforming for hydrogen production) as well as other references [42,43]. As the results (presented in the next section of the paper) show, the proposed SNG plant techno-economic indicators are in line with the published literature sources. Comparing the SNG production using renewable energy with similar fossil fuel alternatives (e.g., coal-to-SNG Great Plains Synfuels Plant, North Dakota, USA), one can notice that the economic values of this analysis were higher than fossil-based options.

4. Results and Discussion

4.1. SNG Technical Performance Indicators

The main goals of the SNG process modeling and simulation were to obtain the mass and energy balances required for the economic evaluation as well as to assess the energy integration elements for the optimization of overall plant energy efficiency. The illustrative SNG plant was designed to be able to process the \( CO_2 \) captured (90% capture rate) from an industrial-size (450 MW) state-of-the-art IGCC power plant. Table 10 shows the main technical performance indicators of a SNG plant that uses electrolytic hydrogen. As can be noted, aside from SNG production (30.8 t/h), the \( CO_2 \) methanation plant also generated a significant amount of electricity (about 28.5 MW net power output). The excess power can be fed into the electric grid or can be used for hydrogen production via the electrolytic process. The power required for hydrogen production and the net efficiency were calculated assuming hydrogen is obtained in a water electrolysis unit. Overall plant energy efficiency was calculated considering the whole system: thermal output of the produced SNG stream, power required for water electrolysis, the power recovered in the steam turbine, and gas decompression steps as well as the ancillary consumption of the plant sub-systems (e.g., pumps, compressors, etc.). The overall results were that the SNG plant showed a high energy efficiency (about 57%).
Table 10. Key technical performance indicators of the proposed SNG plant.

| Main Plant Data                                | U.M. | Value   |
|------------------------------------------------|------|---------|
| Reactant CO\(_2\) flow rate                   | t/h  | 82.51   |
| Reactant H\(_2\) flow rate                    | t/h  | 15.11   |
| Electricity for H\(_2\) production            | kWhe/kg | 53.90 |
| **Total energy input for H\(_2\) production** (A) | MWe  | 814.43  |
| Synthetic natural gas output                  | t/h  | 30.81   |
| Synthetic natural gas LHV                     | MJ/kg | 52.45  |
| SNG thermal output (B)                        | MWh  | 448.88  |
| Power produced by gas decompression           | MWe  | 2.74    |
| Power produced by steam cycle                 | MWe  | 26.95   |
| **Total power production (C)**                | MWe  | 29.69   |
| **Total power consumption (D)**               | MWe  | 1.15    |
| **Net energy efficiency** (B/(A + D − C) × 100) | %    | 57.11   |

SNG decompression is considered to be dependent on the parameters of the gas grid to which the SNG is fed into. The recoverable energy is not included in the energy balance of the plant presented in Table 10. In the economic analysis, however, for the capital cost of this unit, it was considered as the pressure was reduced to 40 bar. This pressure was still significantly higher than the current pressures in common gas grids, but depending on the destination of the SNG, a high pressure could be required.

4.2. SNG Quality Aspects

High CO\(_2\) conversion was targeted to avoid the addition of costly gas separation units. This was achieved using a high pressure and low temperature methanation process. With 99% CO\(_2\) conversion, this can be a satisfactory result for direct use. Inert compound concentration (unreacted CO\(_2\)) in the product stream was around 1% and the remaining unreacted hydrogen was around 4 vol.% as presented in Table 11. Considering a 1:1 blending ratio with natural gas, this would result in a 2 vol.% hydrogen content in the gas grid. This concentration is in line with the European Association of Internal Combustion Engine Manufacturers (EUROMOT) recommendation [27]. The rest of the component’s concentration is presented in Table 11, all of them having low concentrations.

Table 11. Synthetic natural gas flowrate and composition.

| Component | Flowrate (kmole/h) | Vol % |
|-----------|--------------------|-------|
| CH\(_4\)  | 1854.53            | 94.77 |
| CO\(_2\)  | 20.44              | 1.04  |
| CO        | 0.01               | <0.01 |
| H\(_2\)O  | 0.03               | <0.01 |
| H\(_2\)   | 81.80              | 4.18  |
| C\(_2\)H\(_6\) | 0.01       | <0.01 |
| C\(_3\)H\(_8\) | <0.01       | <0.01 |
| Total     | 1956.82            | 100.00|

Based on the obtained concentrations and the description of EUROMOT [27], the resulting SNG would have a WI of around 1% lower than natural gas. This change in WI and calorific value falls within their recommendations for the use of gas in internal combustion engines, accordingly, the SNG can be used within the existing natural gas network.

4.3. SNG Economic Performance Indicators

To achieve the energy independency of a SNG plant, around 300 MW of heat is recovered from the methanation reactor to be used within the process. Accordingly, steam is generated and expanded in a steam turbine for power generation. The turbine has a high impact on the economics of the plant and this is most clear in the case of capital costs. The highest share of the capital cost was the steam cycle, the unit responsible for recovering the
energy generated by the exothermic methanation reaction, as represented in Figure 4. In this case, the power generated was considered to be exported to the grid. The steam cycle was around half of the total capital cost required for the plant. Additional heat exchangers used in the cooling process of the products and in the condensation of the resulting water vapor had the second highest cost. The methanation reactor cost almost 20% of the TIC. The total install cost of the SNG plant was a little below 50 M€, as presented in Table 12. With the cost escalations considered, the total overnight cost was 67 M€, resulting a specific investment cost of about 2166 €/kg synthetic natural gas.

For every considered hydrogen source, a base LC of SNG was calculated. As the methanation plant on its own cannot be economically viable, two additional economic cases were evaluated for each renewable hydrogen source to evaluate the benefits of CO₂ revenue. The first case assumed a 20 €/tCO₂ revenue whereas the second one was 100 €/tCO₂.

Cash flow analysis was used to make the comparison between the technologies based on the LC of SNG. All expenses and revenue were taken into consideration as presented in the Methodology section to compare the economic cases. The LC obtained for the SNG was compared to an average natural gas selling price of 6 €/GJ [4]. The option with a fossil hydrogen source resulted in a cost of SNG of 5.37 €/GJ, which is comparable with the present natural gas market prices. The approximatively 3–4 times higher hydrogen cost observed in the case of renewable hydrogen sources (see Table 5) was also evident in the LC of SNG, this being in the range of 18.62–21.74 €/GJ for the four hydrogen sources assuming biomass as a starting material. The configuration using electrolysis as a hydrogen source gave the same LC of SNG as the biomass gasification as the economic parameters were the same in the two models. The model using PVs and electrolysis for hydrogen production
had the highest production cost at 43.56 €/GJ. These results are represented by the blue bars in Figure 5.

![Figure 5. Levelized cost of synthetic natural gas vs. CO\(_2\) price.](image)

With the addition to the economic model of the CO\(_2\) revenue, the LC dropped, however, this improvement was not enough to counteract the three to four times higher hydrogen cost observed. Applying a 20 €/tCO\(_2\) credit lowered the LC of SNG by around 1 €/GJ, while with the 100 €/tCO\(_2\) credit, the LC dropped with 5.1 €/GJ. Both cases showed improvements when compared to the base economic case with no CO\(_2\) credit, however, it was not enough to lower the LC to the 6 €/GJ threshold. For photo fermentation, dark fermentation, biomass gasification, and bio photolysis, the obtained LC in the 20 €/tCO\(_2\) credit was in the range of 17.60–20.72 €/GJ whereas in the case of the 100 €/tCO\(_2\) credit, it was in the range of 13.52–16.64 €/GJ. The LC obtained in the case of the PV electrolysis base economic case was almost eight times higher than the reference cost, making the CO\(_2\) credit benefits almost unnoticeable. Overall, the LC of SNG, even in the most optimistic scenarios, was still high when compared to the present NG prices. Considering the trend of NG prices from the past several years, the evaluated technologies showed little economic benefit. For comparison reasons, for current industrial projects in the field, the SNG levelized cost produced in the coal-to-SNG Great Plains Synfuels Plant, North Dakota, USA was in the range of 7–11 €/GJ [44].

Of all the hydrogen sources evaluated here, only biomass gasification has the potential to fulfil two important criteria: it is a renewable hydrogen source and also has the potential to produce the required amount of hydrogen to run the plant (15.12 tH\(_2\)/h). Thus, in the next section, only this option is considered.

The LC of SNG obtained consisted of the costs and the revenue of building and running the plant during the economic lifetime. The highest share in the costs, by far, was the hydrogen production cost (as presented in Figure 6). The figure presents the optimistic case with a CO\(_2\) revenue of 100 €/t, which explains the high cost drop observed in the CO\(_2\) revenue section.
Sensitivity analysis performed on the biomass gasification option (Figure 7) also showed a great dependency on the hydrogen cost, this having the highest effect on the cost. The second most important component was the electricity credit obtained from the surplus of power in the plant. Figure 7 also presents the effect of variations in capital cost, operational costs, and interest rate. As can be noted, these components have a small effect on the overall economics of the plant. For other investigated renewable hydrogen production routes, similar variations of SNG levelized cost were observed.

Figure 6. Components of the cash flow analysis.

Figure 7. Variations of the levelized cost of SNG (biomass gasification option).

5. Conclusions

This paper presents the techno-economic evaluation of synthetic natural gas production using renewable hydrogen and captured CO₂. The process is modeled and simulated using Aspen Plus® software. Using the results from process simulation, an in-depth techno-economic model was developed incorporating overall energy efficiency, ancillary energy consumption, capital costs, operational and maintenance costs, SNG levelized cost, sensitivity analysis, etc. Five possible renewable hydrogen sources (photo-fermentation, dark fermentation, biomass gasification, bio-photolysis, and PV electrolysis) were compared to two reference cases (natural gas steam reforming and electrolysis) to assess all options from a techno-economic perspective. All possible renewable hydrogen sources showed a high levelized cost of synthetic natural gas that was mainly attributed to the high cost of hydrogen. The levelized cost of synthetic natural gas was in the range of 18.62–21.74 €/GJ for the evaluated cases, which was around 3.0–3.5 times higher than the actual price of natural gas. Two additional options were added to evaluate the effect of a 20 €/tCO₂ and a 100 €/tCO₂ credit on the SNG levelized cost. Both options significantly improved the
economics of the SNG plant, but not enough to close the gap. Considering the quantity of hydrogen required to run the plant of the evaluated processes, the biomass gasification was the only one that could come into consideration, however, its economics still need to be improved. Although PV electrolysis at this moment has the least attractive economic indicators of the evaluated renewable hydrogen sources, with the further developments of more energy efficient electrolyzers and PV panels, this might be a suitable hydrogen source for future CO₂ utilization technologies.

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