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# Methanol synthesis through CO<sub>2</sub> capture and hydrogenation: thermal integration, energy performance and techno-economic assessment

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## ABSTRACT

This work assesses the opportunity of using “green” methanol (MeOH) produced from renewable electricity as a vector for the decarbonization of chemical process industry. We developed a comprehensive process model to simulate all the relevant sections for the conversion of 1.25 t/h of captured CO<sub>2</sub> from a coal-fired power plant to methanol, using “green” hydrogen via water electrolysis. We applied the pinch analysis methodology for an improved thermal management of the integrated carbon capture, electrolysis and methanol synthesis plant. A network of recovery heat exchangers was designed with the pinch analysis methodology, allowing a thermal energy saving of 4.59 MW, with a net reduction of heating and cooling demands by 81% and 47%, respectively, and an improvement of the global efficiency of the plant from 26.74% to 37.22. We followed a bottom-up approach for the techno-economic assessment, defining a confidence range for the main indicators of system economic viability. We assessed their sensitivity to the cost of electricity from seven renewable energy sources and to the option of selling oxygen produced by water electrolysis, in three different cost scenarios (1-optimistic, 2-realistic, 3-pessimistic). The estimated values of the cost of methanol (COM) span from a maximum range of 2624-2706 €/t (in the case of concentrated solar power) to a minimum of 565-647 €/t (hydropower with highest valorization of electrolytic oxygen), hence resulting in line with the future trends of methanol market price (400-800 €/t) in five of the considered configurations. We have finally estimated the levelized cost of methanol (LCOM) ensuring an internal rate of return ranging from 0 to 10% for each of the techno-economic scenarios identified. Assuming 10% as the target, LCOM in case of hydropower as renewable energy source spans from 874 to 1356 €/t, hence close to the future market price of MeOH with margin of improvement (655-1135 €/t) in case of lower costs of electric energy.

**Keywords:** carbon capture and utilization; electrolysis; CO<sub>2</sub> hydrogenation; methanol synthesis; energy performance; techno-economic assessment.

## Highlights

- Comprehensive process model of a large-scale green power-to-methanol system
- Thermal integration saves 4.59 MW reducing the heating demand by 81%
- We assessed the sensitivity on system economic profitability
- The costs of green and fossil methanol compete in 5 of the studied scenarios
- A production cost of methanol of 874 €/t ensures a 10% internal rate of return

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## List of abbreviations

<b>BEC</b>	Bare Erected Cost
<b>CEPCI</b>	Chemical Engineering Plant Cost Index
<b>COM</b>	Cost of methanol
<b>EBIT</b>	Earnings before interest and taxes
<b>EBITDA</b>	Earnings before interest, taxes, depreciation and amortization
<b>EPCC</b>	Engineering, procurement, and construction cost
<b>FCF</b>	Free Cash Flow
<b>HHV</b>	Higher heating value
<b>IRR</b>	Internal rate of return
<b>LCOE</b>	Levelized cost of electricity
<b>LCOM</b>	Levelized cost of methanol
<b>LHV</b>	Lower heating value
<b>MEA</b>	Monoethanolamine
<b>MeOH</b>	Methanol
<b>NPV</b>	Net present value
<b>OPEX</b>	Operating cost
<b>PBP</b>	Payback period
<b>PC</b>	Process Contingencies
<b>PtX</b>	Power-to-X
<b>RES</b>	Renewable Energy Sources

<b>SPECCA</b>	Specific plant energy consumption for CO <sub>2</sub> avoided
<b>TIC</b>	Total Invested Cost
<b>TOC</b>	Total Overnight Cost
<b>TPC</b>	Total Plant Cost

### List of symbols

<b><math>A</math></b>	-	Amortization of the invested capital
<b><math>B_{1-2}</math></b>	-	Constants for bare module cost (heat exchangers, process vessels and pumps)
<b><math>C_{BM,main}</math></b>	€	Bare module cost of main components
<b><math>C_{BM,tot}</math></b>	€	Total bare module cost
<b><math>C_{BM,WE}</math></b>	€	Total bare module cost of the electrolyzer
<b><math>\tilde{c}_{BM,WE}</math></b>	€/kW	Total specific bare module cost of the electrolysis module
<b><math>c_p</math></b>	kJ/kg/K	Specific heat capacity
<b><math>C_p^0</math></b>	€	Purchased cost of a component (carbon steel, p = 1 bar)
<b><math>E_{elCO_2}</math></b>	MWh/y	Annual electric consumption of the carbon capture section
<b><math>E_{el,sys}</math></b>	MWh/y	Annual electric consumption of the system
<b><math>F_{BM}</math></b>	-	Bare module factor
<b><math>F_M</math></b>	-	Material factor
<b><math>F_P</math></b>	-	Pressure factor
<b><math>i</math></b>	-	Discount rate
<b><math>k</math></b>	year	Years since the investment
<b><math>K_{1-3}</math></b>	-	Parameters of the cost function for the estimation of $C_p^0$
<b><math>\dot{m}</math></b>	kg/s	Mass flow rate
<b><math>m_{CO_2}</math></b>	kt/y	Annual amount of carbon dioxide captured and converted
<b><math>m_{MeOH}</math></b>	t/y	Annual production of methanol
<b><math>n</math></b>	-	Cost exponent
<b><math>n_{life}</math></b>	years	Plant lifetime (25 years)
<b><math>OC</math></b>	€/y	Operating cost in annual cash flow
<b><math>p</math></b>	bar	Absolute pressure
<b><math>P_{WE}</math></b>	kW	Nominal size of the electrolysis stack
<b><math>\dot{Q}_{cooling}</math></b>	kW	Demand of external cooling energy
<b><math>\dot{Q}_{heating}</math></b>	kW	Demand of external heating energy
<b><math>q_{reb}</math></b>	kJ/gCO <sub>2</sub>	Specific regeneration heat duty of the solvent in carbon capture section

$Q_{th_{CO_2}}$	$MWh/y$	Annual thermal consumption of the carbon capture section
$Q_{th_{sys}}$	$MWh/y$	Annual thermal consumption of the system
$R$	€	Revenue in the annual cash flow
$S$	<i>variable</i>	Characteristic size of a component
$T$	°C	Temperature
$Tax$	€	Corporate income tax rate
$V_{H_2O}$	$km^3/y$	Annual water consumption

### Greek symbols

$\alpha$	-	Solvent CO <sub>2</sub> loading
$\Delta T_{min}$	°C	Minimum temperature difference between hot and cold fluids in pinch analysis
$\eta_{gl}$	-	Global efficiency of the Power-to-MeOH system
$\psi$	-	CO <sub>2</sub> removal rate

## 1. Introduction

Since the first Industrial Revolution, fossil fuels have been the main source of primary energy in several sectors of economy such as power plants and industry and they will still probably maintain an important role in the current century [1]. However, the increasing and accelerating emission of greenhouses gases such as carbon dioxide (CO<sub>2</sub>) from the process of combustion [2] has already broken the equilibrium between natural sinks and sources in the so-called “carbon cycle”. Global warming and climate change concerns have recently furthered political efforts to avoid the worst effects of this excess of anthropogenic CO<sub>2</sub> in the atmosphere. Signed in 2015 by 196 countries, the Paris Agreement [3] aims to hold global temperature well below 2 °C above pre-industrial average level, increasing the ability to adapt to the adverse impacts of climate change. To this end, carbon emissions to the atmosphere must peak as soon as possible so that they can decrease later until achieving a neutral balance. In a recent report [4], the Intergovernmental Panel on Climate Change states that limiting global warming to 1.5 °C can be implemented through different mitigation measures: lowering energy and resource intensity, increasing the rate of decarbonization and promoting carbon capture processes.

Carbon capture would be an interim strategy for the stabilization of the emissions due to the use of fossil fuels, but also a long-term solution for the reuse of carbon. In this context, captured CO<sub>2</sub> could be exploited as a raw molecule for the long-term storage of power from RES (renewable energy sources) into chemical form following the Power-to-X (PtX) scheme [5] through reaction with hydrogen produced via different processes, such as water electrolysis. The yielded synthetic gas can be upgraded to many “X” gaseous/liquid fuels and chemicals characterized by high

market value and dispatchability [6]. In particular, future economy will strongly rely on liquid fuels/chemicals due to their capability of storing and displacing remarkable amounts of energy thanks to their high energy density [7].

Methanol (MeOH) is a promising liquid energy carrier with potential use in several applications, either as a chemical or a fuel or as a platform molecule for the synthesis of heavier alcohols, dimethyl ether, gasoline and more complex chemicals, such as olefins [8]. Synthesizing methanol from captured CO<sub>2</sub> and green hydrogen is a valuable opportunity for the decarbonization of chemical process industry and economic sector currently relying on fossil methanol. Table 1 reports the main thermodynamic properties of methanol.

*Table 1 - Physical properties of methanol*

<b>Molecular weight (g/mol)</b>	32.04	<b>Latent heat of vaporization (25 °C, kJ/mol)</b>	37.43
<b>Critical temperature (°C)</b>	239	<b>Vapor pressure (25 °C, bar)</b>	0.1696
<b>Critical pressure (bar)</b>	79.5	<b>LHV (25°C and 1.013 bar, kJ/mol)</b>	638.1
<b>Freezing point (1.013 bar, °C)</b>	-97.6	<b>HHV (25°C and 1.013 bar, kJ/mol)</b>	726.1
<b>Boiling point (1.013 bar, °C)</b>	64.6	<b>Auto-ignition temperature (°C)</b>	470

With a global demand of 60-70 Mt/year in 2015 expected to increase to 190 Mt/year by 2030 [9], more than 80% of methanol production is based on steam reforming of natural gas. However, this process is associated with huge greenhouse gas emissions [10]. The synergy of electrochemical synthesis of hydrogen and its thermocatalytic reaction with captured CO<sub>2</sub> provides alternative pathways for the production of synthetic methanol [11], such as a 1-step direct hydrogenation of CO<sub>2</sub> or a reverse water-gas shift reaction followed by methanol synthesis (as in the case of the CAMERE process, [12]). The overall reduction of greenhouse gas emissions of these pathways with respect to its fossil counterpart is effective only if energy supplied and chemical reactants are characterized by low carbon intensity. Hence, the market for conversion of CO<sub>2</sub> into “green” methanol is strictly driven by the diffusion and storage of RES in chemical form [9]. In fact, the access to clean energy supply and low-cost renewable hydrogen is the main barrier to the commercialization of CO<sub>2</sub>-to-MeOH processes, together with the low efficiency of available catalysts [9,13]; however, pilot programs are in place to investigate these processes and mitigate the associated techno-economic risks [9].

For instance, the “Liquid Sunshine” project in China spans over 50 years and progressively explores the vision of transporting RES throughout the world in the form of methanol [14,15]. Starting from the demonstration of technologies employing hard coal as a raw feedstock of carbon, the project will gradually introduce “sunshine” for electricity supply through PV panels and biomass as carbon source to make the storage and dispatchment chain

fully sustainable and cost-effective [16]. Concluded in 2019, the European MefCO<sub>2</sub> project [17] investigated the techno-economic feasibility of converting renewable H<sub>2</sub> and captured CO<sub>2</sub> to methanol and demonstrated it with a 7.2-t/day CO<sub>2</sub> capture and conversion plant in a coal-fired power station in Germany [18,19]. In Iceland, a commercial-size CO<sub>2</sub>-to-MeOH plant has been operating since 2012 producing 4 kt/y of methanol [20,21]. CO<sub>2</sub> is extracted as a co-product together with steam in a geothermal power plant and captured for reaction in a heterogeneous catalytic reactor with H<sub>2</sub> produced via water electrolysis fed by low-carbon grid electricity [22].

Pushed by the increasing momentum of methanol option for Power-to-Fuels storage, a wide range of literature has recently focused on the study of Power-to-MeOH systems to meet the urgencies of RES diffusion and multi-sectorial decarbonization of economy. Models were developed to simulate the operation of the whole Power-to-MeOH system [10,11,23,24] and to assess thoroughly the techno-economic feasibility [25–27] and the emission savings [28] of the process in different operational scenarios. However, although the sustainability of the process has been discussed from different points of view, little attention has been paid to the potential benefit of heat integration of the different sections of a Power-to-MeOH system on energy performance as well as to the sensitivity of the cost of MeOH to the main economic parameters, such as the electric energy price and the contingencies in the definition of the invested capital.

This work presents a comprehensive model of a full Power-to-MeOH system for the industrial production of “green” methanol through direct hydrogenation of CO<sub>2</sub> captured from a coal-fired power plant (Section 2). Carbon capture system (Section 3.1.1) through amine scrubbing was first modelled starting from real experimental data, validated using the results of the pilot plant designed by Notz et al. [29] as reference and then scaled up to treat a larger flue gas volume. The methanol synthesis (Section 3.1.3) through reaction with hydrogen produced by an alkaline electrolyzer (Section 3.1.2) was modelled taking the cue from the model developed in [30]. In Section 3.2 we highlight the importance of thermal integration of the different sections of the process, so to reduce the thermal needs of the plant. The consumed streams of energy and raw materials resulting from the simulations (Section 4.1) were used to assess the main energy and environmental performance indicators of the Power-to-MeOH system (Section 4.2). Adopting a bottom-up estimation of the total overnight cost (TOC) described in Section 3.3, we computed the levelized cost of MeOH in different scenarios of power supply, providing a confidence range according to reasonable hypotheses on TOC definition and its sensitivity to electric energy price (Section 4.3). Finally, we determined the main energy and economic key performance indicators of the modelled Power-to-MeOH system to assess its viability and prove its economic competitiveness as an option for the decarbonization of methanol industry and market in the considered scenarios.

## 2. Layout of the plant and case-study

Figure 1 presents the scheme of the studied Power-to-MeOH system and its subdivision in three sections:

- Water splitting into hydrogen and oxygen through alkaline electrolyzers;
- Carbon capture with amine-based liquid solvents;
- Synthesis and purification of methanol through direct hydrogenation of captured CO<sub>2</sub>.

As a sustainable alternative to the well-established process of steam methane reforming, we studied the opportunity of using “green” methanol produced from RES as a value-added feedstock for a fully decarbonized chemical process industry. To this end, water electrolysis is fed by energy certified as produced from renewable sources, such as the production of a renewable energy cluster, grid curtailment (i.e. RES overproduction) or low-carbon electric energy purchased from the grid in locations with a very high penetration of RES. The input of renewable electricity is assumed to be constantly available for water electrolysis; the coupling of the system with a real RES availability profile would require also a system of buffers for the raw gases (H<sub>2</sub> and, possibly, CO<sub>2</sub>) suitably sized through a time-resolved model. This is beyond the scope of this paper, in which we developed a model for the nominal operation of the integrated Power-to-MeOH scheme to assess its feasibility from energy and economic standpoints. The study focuses on the production of methanol at a commercial scale (1 m<sup>3</sup>/h of methanol, i.e. about 800 kg/h) and the size of water electrolysis and carbon capture sections was selected in order to feed the methanol synthesis section with sufficient flows of hydrogen and CO<sub>2</sub> captured. We set the size of water electrolysis section equal to 10 MW, hence producing a constant flow rate of 1.9 kNm<sup>3</sup>/h of hydrogen (specific consumption of 5.4 kWh/Nm<sup>3</sup><sub>H<sub>2</sub></sub>) at a pressure of 10 bar. Pure electrolytic hydrogen (99.7%<sub>v</sub> without auxiliary purification equipment) is then compressed for the synthesis of methanol; oxygen is either vented in the base scenario or sold.

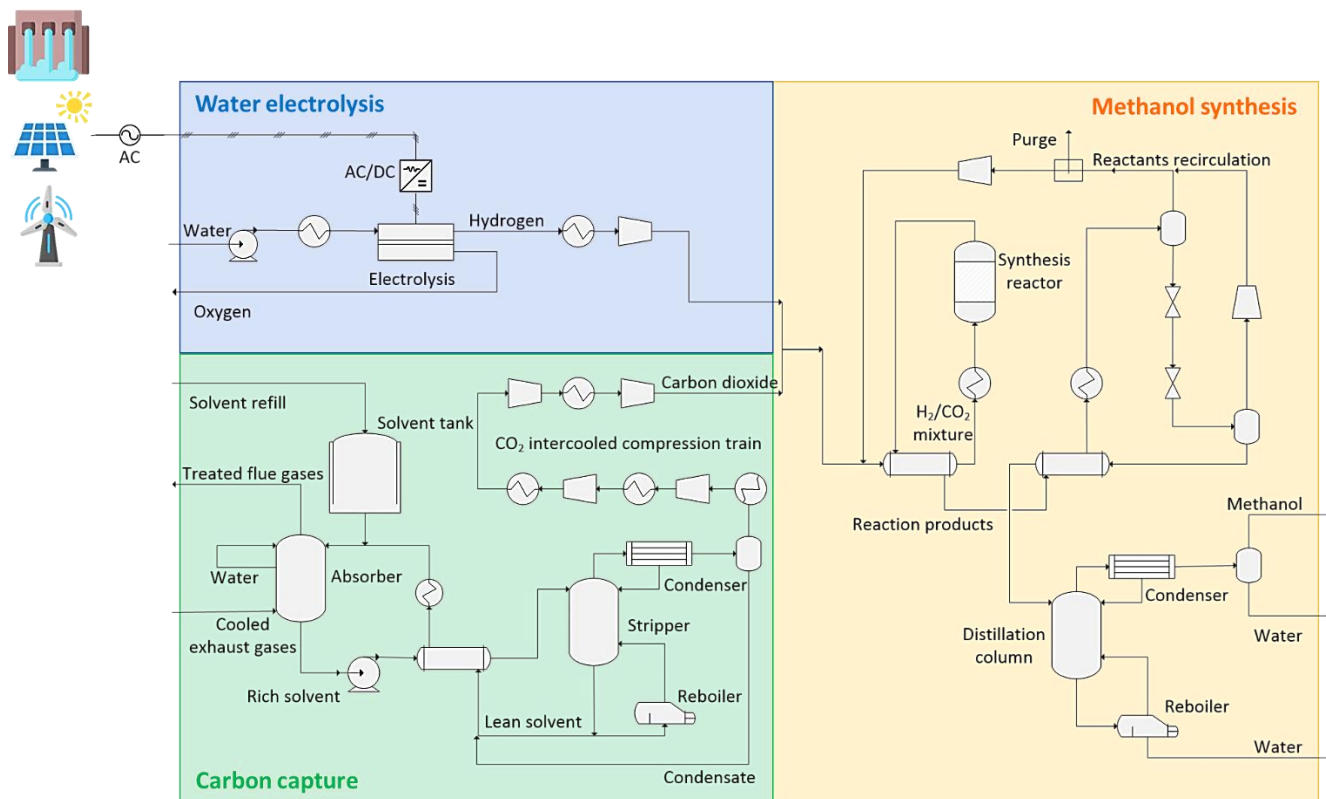


Figure 1 - Layout of the Power-to-MeOH system

In the carbon capture section, a traditional two-column system captures  $632 \text{ Nm}^3/\text{h}$  of  $\text{CO}_2$  (99.7%<sub>w</sub> purity) from  $7.9 \text{ kNm}^3/\text{h}$  of exhausts from a coal-fired power plant through an amine scrubbing process with an aqueous solution of monoethanolamine (MEA), considered as the most mature technology available in the market. The choice of a coal-fired plant as the source of  $\text{CO}_2$  for methanol synthesis is due to the availability of real data of the pilot plant for carbon capture from the exhausts of coal combustion reported in Ref. [29] for the steady-state model of this section. Furthermore, burning hard coal produces exhausts with an intermediate composition between highly concentrated sources of  $\text{CO}_2$  (e.g., biogas upgrading) and diluted point-like streams (e.g., exhausts in gas-fired power plant), hence representing an average source of carbon. Moreover, the elemental composition of exhausts from coal combustion is very close to that of the exhausts from the incineration of municipal solid wastes, an interesting source of  $\text{CO}_2$  for applications in the framework of circular economy.  $\text{CO}_2$  absorption occurs in the absorption column where the solvent flows downwards, in counter-current with the gas mixture coming from the bottom: the  $\text{CO}_2$ -poor gas mixture leaves the top of the absorber. The  $\text{CO}_2$ -rich solvent leaving the absorber from the bottom is pumped and heated up and flows downwards in the desorption column: the solvent is thermally regenerated by a reboiler, so  $\text{CO}_2$  is stripped and leaves the column top. The lean solvent recirculates back to the

absorber through the recovery heat exchanger with a refill in case of leakages for entrainment in the columns and evaporation.

This layout is the most adopted for the capture of CO<sub>2</sub> with liquid solvents through physisorption (e.g., pressurized water scrubbing) and/or chemisorption. Amine scrubbing with MEA represents the state-of-art of chemisorption of CO<sub>2</sub>. The high specific solvent circulation rate and the energy penalty for solvent regeneration make this solution a technologic baseline for carbon capture. Research is currently investigating alternative techniques such as blending or even replacing amines with conventional or functionalized ionic liquids [31] to reduce regeneration heat duty and improve reaction kinetics, calcium looping for high-temperature processes [32], low-temperature adsorbents (e.g., zeolites, MOFs, activated carbons etc.) [33], just to name a few. These promising routes endeavor to raise the energy performance and reduce the costs of carbon capture with respect to the state-of-art technology that is studied in this work.

Captured carbon dioxide flows through an intercooled compression train and mixes with electrolytic hydrogen from water electrolysis section. The pressurized H<sub>2</sub>/CO<sub>2</sub> feed (65 bar) is heated up to 250 °C and fed to the reactor for methanol synthesis through direct hydrogenation of CO<sub>2</sub>: the state of art consists of a reactor equipped with copper-based catalyst – *CuO/ZnO/Al<sub>2</sub>O<sub>3</sub>* (CZA) – operating in the pressure range of 50-100 bar [34,35].

Concerning the catalyst, the use of CZA is the most common option for CO<sub>2</sub> hydrogenation as well as for the traditional methanol synthesis from CO-rich syngas, as this type of catalyst is very active for both processes. The choice of the operating pressure is due to the Cu-based catalyst, which requires a pressure range of 5-10 MPa to obtain a good methanol selectivity (above 50%) in the presence of CO<sub>2</sub> and H<sub>2</sub>, while at lower pressure the rate of formation of CO by reverse water gas shift becomes higher (1-3 orders of magnitude) than that of methanol and thus the resulting CH<sub>3</sub>OH selectivity is extremely low [36]. Several novel catalyst materials are under development with the aim to obtain high methanol selectivity at lower pressure, thus reducing the complexity (and cost) of the synthesis plant. Liao et al. [36] investigated a Pd-Zn catalyst for CO<sub>2</sub> hydrogenation that at 2 MPa showed a CH<sub>3</sub>OH selectivity of 70% compared to 10% over an industrial Cu catalyst. Other works investigated less precious materials, as NiGa [37] or CoGa catalysts [38]. Other approaches involve the improvement of Cu/ZnO using zeolites to add acidic properties and increase reactivity or their confinement in MOFs [19].

The gas mixture leaving the reactor undergoes some steps of pressure reduction and temperature swing for flash separation of gases other than methanol, then a distillation column separates the methanol and a final mechanical separator raises the purity of the produced stream to 99.9% of CH<sub>3</sub>OH.

### 3. Methodology

This section of the paper describes the methodology developed to model the steady-state operation of the three sections of the system (Section 3.1) and their thermal integration with the pinch analysis approach (Section 3.2) in the considered case study. A thorough description of the plant revenue accounting with the cash flow analysis methodology is provided in Section 3.3, while Section 3.4 presents the indicators of the energy and economic performance of the Power-to-MeOH system. A list with a description of the components in the different sections and their corresponding ID is reported in Table S.1 in the Supplementary Material.

#### *3.1. Steady-state process model*

Mass and energy balances and thermodynamic transformations of the operating streams in each of the three sections in Figure 1 were developed using the commercial process simulator Aspen Plus™. As general assumptions for all the modeled sections, 75% isentropic efficiency was set for each compression stage and internal pressure drops were neglected for the circulation pumps and heat exchangers.

##### *3.1.1. Water electrolysis*

This section models the production of the necessary hydrogen for the downstream synthesis of methanol. Water was fed to an equilibrium reactor, a component available in Aspen Plus™ used to reproduce the constant operation at nominal point of 10.26 MW of the alkaline electrolyzer, the most mature, available and affordable technology for low-temperature water electrolysis at commercial scale [39]-[40]. Tap water at  $p = 1$  atm and  $T = 15^\circ\text{C}$  (stream 101 in Figure 2) was pumped and heated to feed the electrolyzer at  $70^\circ\text{C}$  and 10 bar.

Feedwater was dissociated within the electrolyzer according to the reactions listed in Table 2 in a constant flow rate of  $1.9 \text{ kNm}^3/\text{h}$  of  $\text{H}_2$  (assuming a specific energy consumption for electrolysis of  $5.4 \text{ kWh}/\text{Nm}^3_{\text{H}_2}$  [41]) and  $0.95 \text{ kNm}^3/\text{h}$  of oxygen, split through a separator that simulates in the model the electrolyte interposed between the two electrodes of a real alkaline cell.

Table 2 - Chemical reactions employed in the model of the water electrolysis section

Reaction	Type
1) $2 \text{H}_2\text{O} + 2 \text{e}^- \leftrightarrow \text{H}_2 + 2 \text{OH}^-$	Equilibrium
2) $2 \text{OH}^- \leftrightarrow \text{H}_2\text{O} + \frac{1}{2} \text{O}_2 + 2 \text{e}^-$	Equilibrium
3) $\text{H}_2\text{O} \leftrightarrow \frac{1}{2} \text{O}_2 + \text{H}_2$ (overall)	Equilibrium

Excess water at the outlet was recirculated back to the reactor (i.e., to the stack inlet). Hydrogen was produced (stream 105) at high purity (99.7%) and pure oxygen (stream 104) is vented as represented in Figure 2; however, it can be sold in order to increase the revenues of the plant, as explained in Section 4.3.4.

Since alkaline electrolysis is an exothermic process working over thermoneutral voltage for a higher productivity, a cooling system is designed to remove the heat generated at nominal power with the purpose of maintaining the nominal operating conditions (Table S.2 in Supplementary Material).

Finally, the produced hydrogen was cooled and compressed (stream 107) to the operating conditions required at the inlet of the methanol synthesis section.

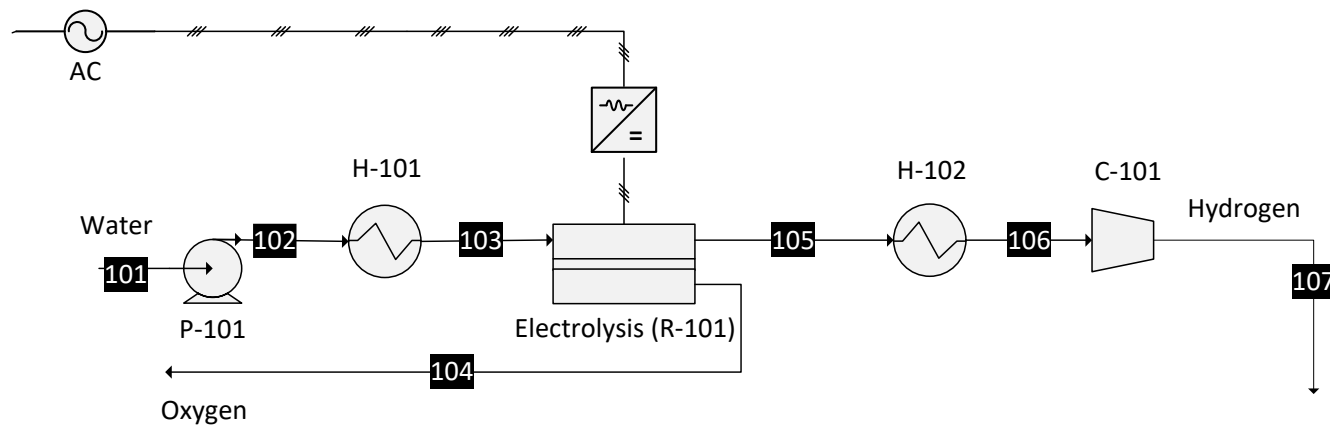


Figure 2 - Flowsheet of the model of water electrolysis section. Recirculation of excess feedwater is not shown

### 3.1.2. Carbon capture through amine scrubbing

The process of carbon capture from the exhaust gas of a coal-fired power plant through amine scrubbing was developed taking the cue from the real datasheet and operating results of the pilot plant described in Ref. [29]: Figure 3 shows the flowsheet of the model reporting the tags for the operating streams and the main blocks.



Table 3 - Chemical reactions employed in the model of the carbon capture section

Reaction	Type	Reaction	Type
4) $\text{H}_2\text{O} + \text{HCO}_3^- \leftrightarrow \text{CO}_3^{2-} + \text{H}_3\text{O}^+$	Equilibrium	9) $\text{OH}^- + \text{CO}_2 \rightarrow \text{HCO}_3^-$	Kinetic
5) $2 \text{H}_2\text{O} \leftrightarrow \text{OH}^- + \text{H}_3\text{O}^+$	Equilibrium	10) $\text{HCO}_3^- \rightarrow \text{OH}^- + \text{CO}_2$	Kinetic
6) $2 \text{H}_2\text{O} + \text{CO}_2 \leftrightarrow \text{HCO}_3^- + \text{H}_3\text{O}^+$	Equilibrium	11) $\text{MEA} + \text{CO}_2 + \text{H}_2\text{O} \rightarrow \text{MEACOO}^- + \text{H}_3\text{O}^+$	Kinetic
7) $\text{MEACOO}^- + \text{H}_2\text{O} \leftrightarrow \text{MEA} + \text{HCO}_3^-$	Equilibrium	12) $\text{MEACOO}^- + \text{H}_3\text{O}^+ \rightarrow \text{MEA} + \text{CO}_2 + \text{H}_2\text{O}$	Kinetic
8) $\text{MEAH}^+ + \text{H}_2\text{O} \leftrightarrow \text{MEA} + \text{H}_3\text{O}^+$	Equilibrium		

The thermodynamic properties of the chemical species involved in the carbon capture process were estimated with the unsymmetrical electrolyte non-random two-liquid method for the liquid phase, Redlich-Kwong equation and Henry's Law for gas phase. In two-phase blocks like the ab-/desorption columns, the mass transfer was modelled with a "two-film" rate-based approach neglecting the mixing by convection [43,44].

The developed model was tuned in order to follow the specifications and the results reported by Notz et al. [29]. It is worth remarking that the carbon capture loop was modelled as an open system in order to avoid convergence issues with Aspen Plus™: the lean solvent regenerated in the stripper (stream 216) is not fed directly to the absorption column. Instead, we employed a fictitious feed of lean solvent (stream 206) characterized by the same loading  $\alpha$  reported in the consulted experimental results [29]:

$$\alpha = \frac{\text{moles of } \text{CO}_2 \text{ species}}{\text{moles of MEA species}} = 30.8\% \quad (\text{Eq. 1})$$

The non-zero loading of the solvent fed to the absorber accounts for the fact that the solvent cannot be completely regenerated in a real closed-loop continuous operation (i.e. a part of  $\text{CO}_2$  keeps being dissolved in or chemically bound to the solvent) unless a high amount of heat is consumed in the reboiler of the stripper. To ensure the mass balance of the lean solvent, we introduced a design specification that determines the regeneration heat duty in the stripper reboiler (H-201) needed to guarantee that the loading of stream 216 equals the loading of stream 206.

The mass streams and the energy fluxes of the developed model were compared with those of the reference pilot plant and the model was validated considering the following operating parameters:

$$\text{CO}_2 \text{ removal rate, } \Psi = \frac{\dot{m}_{218}(\text{CO}_2)}{\dot{m}_{201}(\text{CO}_2)} \quad (\text{Eq. 2})$$

$$\text{Specific regeneration heat duty, } q_{reb} = \frac{\dot{Q}_{H-201}}{\dot{m}_{218(CO_2)}} \quad (\text{Eq. 3})$$

$$\text{Liquid solvent to flue gas ratio in the absorber, } \frac{L}{G} = \frac{\dot{m}_{206}}{\dot{m}_{201}} \quad (\text{Eq. 4})$$

Where  $\dot{m}_{218(CO_2)}$  and  $\dot{m}_{201(CO_2)}$  (kg/s) are the mass flow rates of CO<sub>2</sub> in streams 218 and 201, respectively;  $\dot{Q}_{H-201}$  (kW) is the regeneration heat duty consumed by the stripper reboiler;  $\dot{m}_{206}$  and  $\dot{m}_{201}$  are the mass flow rates of streams 206 and 201, respectively. The following table reports the results of the validation of the developed model.

Table 4 - Validation of the developed model

Operating parameter	Results of the pilot plant [29]	Developed model	Deviation
CO <sub>2</sub> removal rate, $\Psi$	51.3%	51.7%	0.78%
Specific regeneration heat duty, $q_{reb}$ (kJ/gCO <sub>2</sub> )	4.68	4.64	0.85%
L/G ratio	2.76	2.76	-

The validated model was then scaled up to the industrial size of the studied Power-to-MeOH system, keeping the operating conditions, the design specifications and the configurations of each component unchanged, apart the size of the absorption and desorption columns (Table S.4 in Supplementary Material). The typical value of  $\Psi$  used for the analysis of modelled carbon capture process [45–47] and the design of real amine scrubbing plants [48,49] is 90%, which is higher than the value from the reference plant used in the validated model (51.7%). Hence, we adopted a CO<sub>2</sub> removal rate of 75% for the scaled-up model of the carbon capture section as a trade-off between the mentioned data, also considering that Notz et al. [29] report another experiment at lower CO<sub>2</sub> partial pressure resulting with  $\Psi = 75\%$ . From this point on, the analysis refers uniquely to the validated scaled-up model.

Table S.3 in the Supplementary Material reports the values of the chemical and physical properties of the streams entering the carbon capture section, i.e. flue gas mixture coming from the upstream power plant (stream 201 in Figure 3), the fresh solvent refilled in the ab-/desorption loop in order to compensate the leakages due to entrainment and evaporation (stream 226 in Figure 3) and the fictitious lean solvent stream 206.

Table S.4 in Supplementary Material lists the detailed specifications of the two ab-/desorption columns: these blocks were simulated as rate-based multi-stage vapor-liquid fractionating unit (RadFrac components in Aspen Plus™) with two-film approximation.

### 3.1.3. Methanol synthesis

In this section, the captured carbon dioxide is hydrogenated and converted into methanol. In order to simulate this process, the model developed by Calogero [30] and based on the work by Atsonios et al. [25] was used as reference. The model was rescaled to fit the flow rate of CO<sub>2</sub> captured in our case study and integrated in our model.

As shown in Figure 4, the feeding stream 301 (characterized by an H<sub>2</sub>/CO<sub>2</sub> ratio equal to 3), was mixed with the recirculated excess reactants (stream 311). In order to guarantee the operating conditions of T=250°C and p=65 bar (see Table S.5 in Supplementary Material), the feeding mixture is preheated before entering the reactor (recovery heat exchanger E-301 and heater H-301).

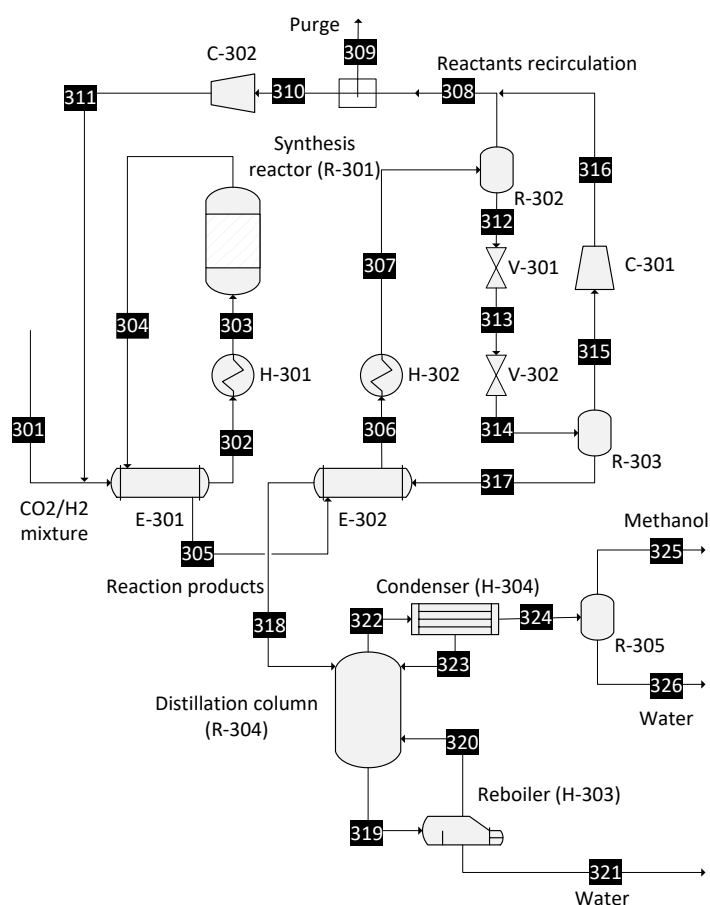


Figure 4 - Flowsheet of the model of methanol synthesis section

The reactor for methanol synthesis (see Table S.6 in Supplementary Material for the specifications) was modelled as a horizontal isothermal plug-flow reactor with the kinetic scheme proposed by Calogero [30]. The Cu/ZnO/Al<sub>2</sub>O<sub>3</sub> solid catalyst in the reactor enhanced the hydrogenation of CO<sub>2</sub>, producing methanol according to the reactions reported in Table 5 proposed by Graaf's kinetic model [50].

Table 5- Chemical reactions employed in the model of the methanol synthesis section

Reaction	Enthalpy Variation
13) $\text{CO}_2 + 3 \text{H}_2 \leftrightarrow \text{CH}_3\text{OH} + \text{H}_2\text{O}$	$\Delta H_{25^\circ\text{C}} = -49.5 \text{ kJ/mol}_{\text{CO}_2}$
14) $\text{CO}_2 + \text{H}_2 \leftrightarrow \text{CO} + \text{H}_2\text{O}$	$\Delta H_{25^\circ\text{C}} = +41.2 \text{ kJ/mol}_{\text{CO}_2}$
15) $\text{CO} + 2 \text{H}_2 \leftrightarrow \text{CH}_3\text{OH}$	$\Delta H_{25^\circ\text{C}} = -90.7 \text{ kJ/mol}_{\text{CO}_2}$

After the reactions, the pressure and the temperature of the fluid were decreased through two throttling valves and a recovery heat exchanger in order to reach the thermodynamic conditions required in the distillation column ( $T=85^\circ\text{C}$ ,  $p=2.2 \text{ bar}$ ). The distillation column was modeled considering 60 stages with reboiler pressure of 1.1 bar and a pressure drop of 0.1 bar along the column height. A mechanical demister was added to separate water from the distillate in order to obtain methanol with a degree of purity of 99.9% (stream 325).

### 3.2. Thermal integration

The demand of external energy in terms of heating ( $\dot{Q}_{heating}$ ) and cooling needs ( $\dot{Q}_{cooling}$ ) of the whole Power-to-MeOH was minimized exploiting the endothermicity and exothermicity of the different components. To this end, the pinch analysis methodology was applied to identify the hot and cold fluids (Figure 5) and to design a network of recovery heat exchangers for an improved energy integration of the plant. An arbitrary minimum temperature difference ( $\Delta T_{min}$ ) of  $10^\circ\text{C}$  was assumed for the coherent coupling of different streams, while an average specific heat capacity ( $c_p$ ,  $\text{kJ/kg/K}$ ) was set in the range between inlet and outlet temperatures of each stream.

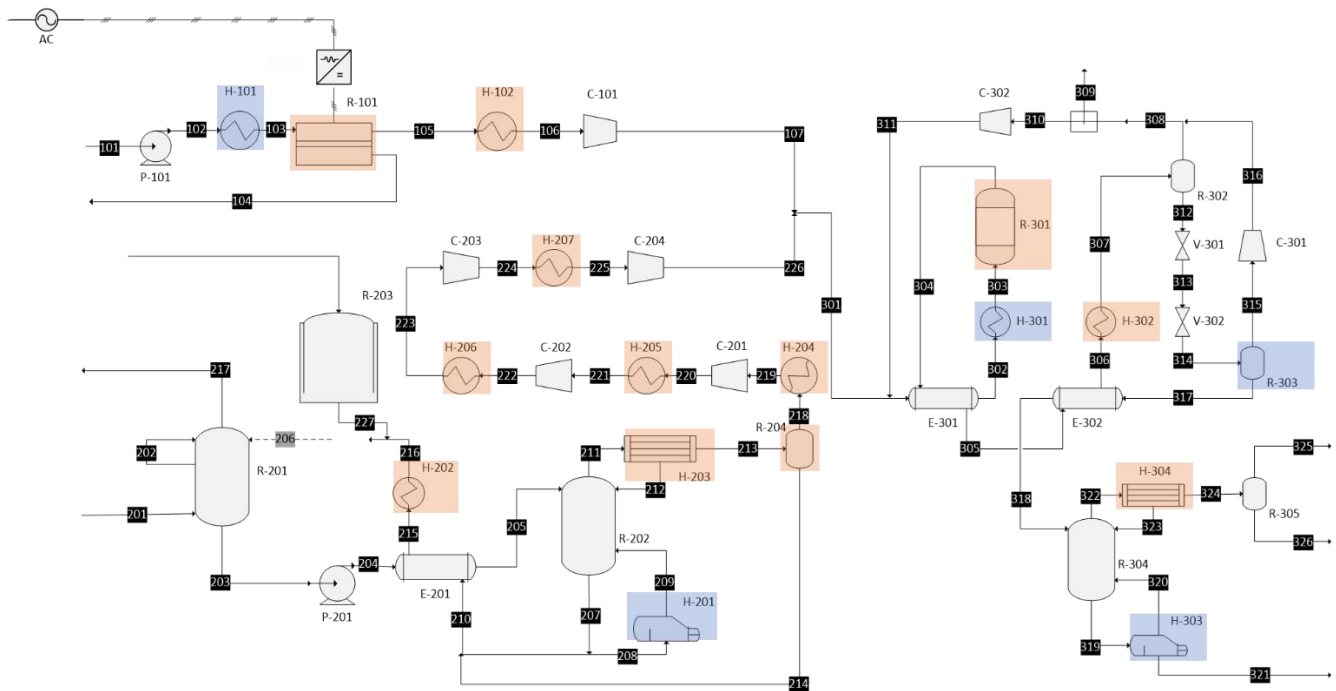


Figure 5 - Identification of the hot fluids (red boxes) and cold fluids (blue boxes) involved in the pinch analysis

### 3.3. Techno-economic assessment

The results of the steady-state models of the three sections and their thermal integration were used as input data for the assessment of the techno-economic feasibility of the Power-to-MeOH system through the revenue accounting methodology. The boundaries of the techno-economic assessment include the components represented in the colored boxes in Figure 1 and the streams and energy fluxes entering and leaving the whole system. For the analysis, we introduced three different techno-economic scenarios in order to provide a -15%/+30% confidence range in which the values of the TOC and the main economic indicators are expected to be:

- Scenario 1 – optimistic, with economic assumptions to define the lower value of the TOC;
- Scenario 2 – realistic, relying on data from pilot plant projects, in literature and from reports of R&D organizations [51–54].
- Scenario 3 – pessimistic, corresponding to the upper bound of the confidence range for TOC.

In all the scenarios, we assumed a project lifetime of 25 years and 8000 operating hours per year.

### 3.3.1. CAPEX estimation: scenarios and assumptions

A bottom-up approach was adopted for the definition of the TOC, employing the sizes ( $S$ ) of the different components in the plant as input variables of cost functions available in literature. For each of the main components present in the studied system (packed columns, reactors, tanks, heat exchangers, pumps, compressors and flash separators), the cost functions from Turton et al. [52] were employed to evaluate the purchased cost  $C_p^0$  assuming carbon steel as construction material and operating pressure of 1 bar:

$$\log_{10} C_p^0 = K_1 + K_2 \log_{10} S + K_3 [\log_{10} S]^2 \quad (\text{Eq. 5})$$

The parameters  $K_1$ - $K_3$  and the range of sizes  $[S_{min}, S_{max}]$  of validity of the cost functions were also provided in Ref. [52]. If the size  $S$  of the component was out of the  $[S_{min}, S_{max}]$  range, we applied the following approximation with the cost exponent  $n$  equal to 0.6 (six-tenths rule) in lack of further indications [55]:

$$C_p^0 = \begin{cases} C_p^0(S_{min}) \cdot \left(\frac{S}{S_{min}}\right)^n, & S < S_{min} \\ C_p^0(S_{max}) \cdot \left(\frac{S}{S_{max}}\right)^n, & S > S_{max} \end{cases} \quad (\text{Eq. 6})$$

In order to estimate the bare module cost of the effective component, one should introduce the factor  $F_M$  accounting for the real construction material and the factor  $F_P$  accounting for changes in operating pressure:

$$C_{BM} = C_p^0 \cdot F_{BM} = C_p^0 \cdot (B_1 + B_2 F_P F_M) \quad (\text{Eq. 7})$$

Turton et al. [52] report the factor  $F_M$  according to the considered construction material and the constants  $B_1$  and  $B_2$  for heat exchangers, vertical/horizontal process vessels and pumps; for the remaining components, the bare module factor  $F_{BM}$  is provided. The cost functions provided by Turton et al. refer to the year 2001; we updated the bare module costs to year 2019 using the ratio of CEPCI (Chemical Engineering Plant Cost Index, [56]) values to account for the effect of inflation:

$$C_{BM}(2019) = C_{BM}(2001) \cdot \frac{CEPCI_{2019}}{CEPCI_{2001}} = C_{BM}(2001) \cdot \frac{618,7}{394} \quad (\text{Eq. 8})$$

The first value of the \$/€ exchange rate in 2019 (1 € = 1.145 \$) was used for conversion from US dollars (the reference currency used in [52]) to Euros.

Once determined  $C_{BM,main}$  as the sum of all the bare module costs of the main components, the Bare Erected Cost (BEC) of the plant is estimated considering the cost of minor equipment and machinery and the supporting work. These items are estimated as a fraction of  $C_{BM,main}$ , as explained in the following table:

Table 6 - Scenarios and assumptions for the estimation of the BEC of the plant [52]

	Sc.1 - optimistic	Sc.2 - realistic	Sc.3 - pessimistic
$C_{BM,main}$	71.2%	68%	63.1%
<b>Pipe, valves and fittings</b>	18.0%	20.0%	23.0%
<b>Process instruments &amp; control</b>	6.3%	7.0%	8.1%
<b>Electric equipment and materials</b>	4.5%	5.0%	5.8%
$C_{BM,tot}$	100%	100%	100%
$C_{BM,tot}$	71.2%	68%	63.2%
<b>Erection and installation labor</b>	21.6%	24%	27.6%
<b>Buildings, materials and labor</b>	7.2%	8%	9.2%
<b>BEC</b>	100%	100%	100%

Concerning the electrolysis system, the following cost function valid for the total cost of alkaline electrolyzers with size bigger than 2 MW (including the stack, power converters, water purification system, gas purification unit and water recirculation pump) was derived fitting the data reported in [57]:

$$\tilde{c}_{BM,WE} = 1437 \cdot P_{WE}^{-0.095} \quad (\text{Eq. 9})$$

where  $\tilde{c}_{BM,WE}$  is the total specific cost of the electrolysis module (€/kW) and  $P_{WE}$  is the nominal stack size (kW). We obtained a specific cost of 599 €/kW for a 10-MW electrolyzer (i.e., the total bare module cost  $C_{BM,WE}$  is 5.99 M€).

We followed the guidelines reported by NETL [51] for the estimation of the TOC, with the assumptions listed in Table 7. Process contingencies compensate for performance uncertainties and are applied as a fraction of BEC to each section of the plant according to the level of maturity of the technology.

Table 7 - Scenarios and assumptions for the estimation of the TOC [51]

	Sc.1 - optimistic	Sc.2 - realistic	Sc.3 - pessimistic
<b>EPC contractor services (%BEC)</b>	15.0%	17.5%	20.0%
<b>Process contingencies PC (%BEC)</b>	*	*	*
<b>Project contingencies %(EPCC+PC)</b>	15.0%	15.0% [54]	30.0%
<b>Start-up cost (%TPC)</b>	1.8%	2.0%	2.3%
<b>Inventory (%TPC)</b>	0.45%	0.50%	0.58%
<b>Financing costs (%TPC)</b>	2.4%	2.7%	3.1%
<b>Other owner's costs (%TPC)</b>	13.5%	15.0%	17.3%

\* PC: 0% – 5% – 10% for water electrolysis section;  
 5% – 17.6% [53] – 20% for carbon capture section;  
 0% – 5% – 10% for methanol synthesis section.

### 3.3.2. OPEX estimation: scenarios and assumptions

For the techno-economic assessment we also considered the following operating costs (OPEX) of the system, with  $OC_{1-7}$  being variable operating costs proportional to the productivity of the Power-to-MeOH system and  $OC_{8,9}$  being fixed costs not dependent on the flows of energy and/or mass consumed by the plant:

- $OC_1$ . Electric energy consumed for the self-consumption of the Balance of Plant (compressors, pumps, etc.) and to feed water electrolysis.
- $OC_2$ . Water supply for the different processes (water electrolysis, solvent makeup in carbon capture section), with a unit cost of 3.36 €/m<sup>3</sup>.
- $OC_3$ . Tap water ( $c_p = 4.186$  kJ/kg/K) operating between 12 °C and 15 °C used to cool down streams that are at temperature higher than or equal to 25 °C, with a unit cost of 3.36 €/m<sup>3</sup>.
- $OC_4$ . Hot utility to fulfil the thermal energy demand of the system through electric heaters with a power-to-heat efficiency of 95%.
- $OC_5$ . Refill of MEA in carbon capture section due to loss of solvent, with a unit cost of 1.09 €/kg [47].
- $OC_6$ . Replacement of the electrolysis stack at 11<sup>th</sup> and 21<sup>st</sup> year of operation with the following cost [58]:

$$OC_6 = \frac{2 \cdot C_{BM,WE} \cdot 0.4}{3} \quad (\text{Eq. 10})$$

- $OC_7$ . Replacement of the Cu/ZnO/Al<sub>2</sub>O<sub>3</sub> catalyst in the methanol synthesis reactor, with a unit cost of 95.24 €/kg [28].
- $OC_8$ . Maintenance, assumed as 3% of Total Plant Cost (TPC) [47].

- OC<sub>9</sub>, Insurance, assumed as 1% of Total Plant Cost (TPC) [47].

Labor costs were excluded from the analysis since they strongly depend on the plant location. All the OPEX items are assumed to be constant during the plant lifetime. Concerning OC<sub>1</sub> and OC<sub>4</sub>, we considered different options to fulfil the electric demand of the whole system with RES. The following table reports the levelized cost of electricity (LCOE) for each of the considered renewable sources in 2018 [59], considering a \$/€ exchange rate of 1.145:

Table 8 - Global electricity costs from RES in 2018 [59]

Renewable source	Scenario	Global weighted-average cost of electricity (€/MWh, 2018)	Cost of electricity: 5 <sup>th</sup> and 95 <sup>th</sup> percentiles (€/MWh, 2018)
Bioenergy	A	54.15	41.92 – 212.23
Geothermal	B	62.88	52.4 – 124.89
Hydro	C	41.05	26.2 – 118.78
Solar PV	D	74.24	50.66 – 191.27
Conc. solar power	E	161.57	95.2 – 237.55
Offshore wind	F	110.92	89.08 – 172.93
Onshore wind	G	48.91	38.43 – 87.34

As explained in Section 3.4, we did not consider the sale of methanol at a market price in the cash flow analysis, rather we computed its production cost (COM, cost of methanol) that allows to reach a breakeven at the end of plant lifespan. Apart from methanol, the only other stream of the Power-to-MeOH system (shown in Figure 1) with an interesting market value is oxygen. We will discuss (Section 4.3.4) the sensitivity of the economic figures of merit (Section 3.4) to the sale of oxygen, considering a range for its market price:

- 0 €/t (i.e., oxygen is vented);
- 70 €/t [60,61] associated to liquefied oxygen;
- 150 €/t [62] associated to oxygen compressed at 30 bar.

### 3.3.3. Cash flow analysis and revenue accounting methodology

The economic feasibility of the investment project was verified with the cash flow analysis methodology, with the assumptions of no discount nor escalation rate when estimating the annual cash flows during the plant lifetime and of no remuneration of the invested capital. The following equations were used, where  $k$  represents the time passed since the investment (year 0):

- Earnings before interest, taxes, depreciation and amortization:

$$EBITDA_k = \sum_i R_i - \left| \sum_j OC_j \right| \quad (\text{Eq. 11})$$

with  $OC_j$  referring to the operating costs listed in Section 3.3.2 and  $R_i$  to the revenue from sale of methanol and oxygen (if considered, as already anticipated in Section 3.3.2 and thoroughly discussed in Section 4.3.4).

- Amortization of the invested capital:

$$A_k = \frac{|TOC|}{n_{life}} \quad (\text{Eq. 12})$$

where  $n_{life}$  is the plant lifetime (25 years)

- Earnings before interest and taxes:

$$EBIT_k = EBITDA_k - |A_k| \quad (\text{Eq. 13})$$

- Taxes:

$$Tax_k = \%_T \cdot EBIT = 24\% EBIT \quad \text{if } EBIT > 0 \quad (\text{Eq. 14})$$

where 24% is the corporate income tax rate in law in Italy since 2017.

- Free Cash Flow:

$$FCF_k = EBITDA_k - |Tax_k| \quad (\text{Eq. 15})$$

- Total Investment Cost, i.e. the total flow of money expended (costs) during the plant lifetime:

$$TIC = -|TOC| - \sum_{k=1}^{n_{life}} \left| \sum_j OC_{j,k} \right| \quad (\text{Eq. 16})$$

### 3.4. Key Performance Indicators of the system

The performance of the system was assessed from both technical and economic standpoints, with particular attention to the reduction of the impacts with respect to the zero-alternative.

- The main environmental benefit of the studied system consists of the reduction of carbon emissions to the atmosphere:  $m_{CO_2}$  indicates the mass (t/y) of carbon dioxide captured and converted to methanol on annual basis.
- Water consumption represents an important issue for the sustainability of the process: hence, we also assessed the annual amount ( $m^3/y$ ) of tap water  $V_{H_2O}$  consumed in the different sections.
- Specific plant energy consumption for  $CO_2$  avoided (SPECCEA, kWh/kg $_{CO_2}$ ), i.e., the total consumption of electric energy and heat (assuming that they have the same exergy level) of carbon capture section to remove a unit of  $CO_2$  from the flue gas stream and convey it to the methanol synthesis section:

$$SPECCEA = \frac{E_{el_{CO_2}} + Q_{th_{CO_2}}}{m_{CO_2}} \quad (\text{Eq. 17})$$

- The global efficiency of the Power-to-MeOH plant, considering the electric ( $E_{el_{sys}}$ ) and thermal consumption ( $Q_{th_{sys}}$ ) of the whole system to produce the annual amount of MeOH ( $m_{MeOH}$ ), characterized by the lower heating value  $LHV_{MeOH}$  reported in Table 1:

$$\eta_{gl} = \frac{m_{MeOH} \cdot LHV_{MeOH}}{E_{el_{sys}} + Q_{th_{sys}}} \quad (\text{Eq. 18})$$

- As main indicator of the techno-economic feasibility of the system, we assessed the cost of methanol (COM, €/kg $_{MeOH}$ )

$$COM = \frac{-TIC - |R_{O_2}|}{m_{MeOH} \cdot n_{life}} \quad (\text{Eq. 19})$$

as the production cost of methanol that ensures a breakeven over the project lifetime of 25 years. We compared it with the current and forecasted market prices of methanol. COM was assessed in the three different scenarios for the definition of the TOC (Section 3.3) and considering the three options for oxygen sale ( $R_{O_2}$ ) presented in Section 3.3.2 (selling price: 0-70-150 €/t). In the most and in the least competitive configurations, we also estimated the values of levelized cost of methanol (LCOM) ensuring an internal rate of return (IRR) of the investment ranging from 0 to 10%:

$$IRR = i: \sum_{k=1}^{n_{life}} \frac{FCF_k}{(1+i)^k} - |TOC| = NPV = 0 \quad (\text{Eq. 20})$$

Since the discount rate strongly depends on investment structure and market conditions and affects the cash flow analysis, we preferred not to assess NPV (net present value), rather to provide the LCOM associated to the parameterized values of the discount rate (i.e., IRR) that ensures a breakeven at the end of plant lifetime – or, in other words,  $NPV = 0$ .

- As an additional indicator, we provided the Payback Period (PBP), i.e., the number of years it takes to break even from the investment considering the annual net free cash flows:

$$PBP = \min \left\{ k \mid -|TOC| + \sum_{k=1}^{n_{life}} FCF_k > 0 \right\} \quad (\text{Eq. 21})$$

## 4. Results and discussion

We modelled the full Power-to-MeOH chain with the capture of about 1.25 t/h of carbon dioxide in 10.2 t/h of exhausts from a coal-fired power plant and its further conversion to 788 kg/h of methanol through a catalytically driven thermochemical reaction with 1.9 kNm<sup>3</sup>/h of electrolytic hydrogen. The outcomes of these simulations are presented in this section, along with the results of thermal integration, energy analysis and economic assessment.

### 4.1. Steady-state simulation of the plant

This section reports the energy balance resulting from the simulations and the results obtained from the thermal integration calculations.

#### 4.1.1. Energy balance

Table 9 summarizes the energy balance of the full Power-to-MeOH chain, reporting the exchanges of power and heat in the different sections of the plant to produce 788 kg/h of methanol (i.e. about 4.36 MW of chemical power, considering the chemical data in Table 1). Considering the most representative streams of water electrolysis, carbon capture and methanol synthesis sections, the relevant information about mass balance and the thermodynamic conditions of each streams resulting from the simulations are reported in Table S.7 in the Supplementary Material. The IDs refer to Figure 5.

The left-hand side of Table 9 reports the heat fluxes exchanged between the plant components and the environment (negative values refer to the cooling duties). The main thermal fluxes exceeding 1 MW derive from the cooling of excess feedwater recirculation in the electrolyzer, the carbon capture solvent heating for CO<sub>2</sub> stripping, the preheating of the H<sub>2</sub>/CO<sub>2</sub> reactant mixture entering the methanol synthesis reactor and the cooling of products mixture at the inlet of distillation column. The presence of many heat fluxes with opposite signs justifies the investigation about thermal integration of the different sections to reduce cooling and heating demands of the plant.

Concerning electric energy consumption on the right-hand side of the table, power demanded to run electrolysis is 2-3 orders of magnitude higher than other contributions listed, which sum up to about 392 kW. In order to certify the produced methanol as “green”, the power demand should be fulfilled by a sufficiently high renewable generation (scenarios A-G in Table 8), hence excluding any indirect emission of CO<sub>2</sub>. We accounted for this renewable energy consumption in the energy balance and the KPIs, without modelling its daily/seasonal fluctuations. A first rough estimation of 42.48% for the power-to-MeOH process gross efficiency is computed considering the power demanded to run electrolysis as the only energy consumed by the plant and converted into the chemical power of 788 kg/h of methanol. This figure of merit will be compared with the values of the global plant efficiency (Section 4.2) accounting also for the thermal needs.

Table 9 – Energy balance of the Power-to-MeOH system

<b>ID</b>	<b>Component Description</b>	<b>Heat duty (kW)</b>	<b>ID</b>	<b>Component description</b>	<b>Electric power (kW)</b>
H-101	Electrolysis water preheating	116.28	P-101	Electrolysis feedwater pumping	0,56
R-101	Rec. excess water cooling	-3482	R-101	Electrolysis module	10260
H-102	Hydrogen cooling	-37.48	C-101	Hydrogen compression	194.92
E-201	Solvent recovery heat exchanger	±1804.7	P-201	Rich solvent pumping	2.76
H-201	Reboiler of the stripper	1268.52	C-201	CO <sub>2</sub> post-compression	25.85
H-202	Lean solvent cooling	-502.4	C-202	CO <sub>2</sub> post-compression	25.52
H-203	Condenser of the stripper	-150.14	C-203	CO <sub>2</sub> post-compression	25.06
H-204	CO <sub>2</sub> intercooling stage #1	-18.36	C-204	CO <sub>2</sub> post-compression	24.68
H-205	CO <sub>2</sub> intercooling stage #2	-26.75	C-301	Compression of the rec. reactants	92.71
H-206	CO <sub>2</sub> intercooling stage #3	-26.66	C-302	Compression of the rec. reactants	≈ 0
H-207	CO <sub>2</sub> intercooling stage #4	-25.68			
E-301	Recovery heat exchanger	±909.94	<b>ID</b>	<b>Component description</b>	<b>Chemical content (kW)</b>
E-302	Recovery heat exchanger	±125.14	326	Methanol production	4358.2
H-301	Reactants preheating	1030.53			
R-303	Flash sep. of MeOH mixture	77.36			
H-302	Cooling of products mixture	-1820.4			
H-303	Reboiler of the distill. column	313.19			
R-301	MeOH synthesis heat of reaction	-426.05			
H-304	Condenser of the distill. column	-426.11			

#### 4.1.2. Results of pinch analysis

As reported earlier, the fulfilment of thermal energy demands for the conditioning of the streams operating in the plant heavily affects the system feasibility. The presence of different fluids heating and cooling in the sections offers the opportunity of reducing the thermal demand of the plant. Hence, we applied pinch analysis methodology to minimize the heat supplied from external thermal sources through a network of recovery heat exchangers.

Setting an arbitrary minimum temperature difference ( $\Delta T_{\min}$ ) of 10 °C, the pinch point ( $T_{pp}$ ) of the plant was found at 108.5°C. Table 10 reports the sensible reduction of both cooling (-47%) and heating (-81%) demand after the thermal integration of the different sections of the plant, with a beneficial effect on the energy performance of the process. Of the 5193 kW of cooling duty after integration, about 4767 kW are associated to streams cooling down at temperatures higher than 25°C: this cooling duty will be fulfilled using tap water undergoing an increase in temperature equal to 3°C. The remaining 8% of cooling demand corresponds to the duty of the condenser in the distillation column operating at around -18°C: considering its limited weight on the global energy demand, this contribution is neglected in the techno-economic assessment and the evaluation of the KPIs of the plant. The heating demand resulting after thermal integration (1057.3 kW) is almost totally associated to the preheating of H<sub>2</sub>/CO<sub>2</sub> reactant mixture (cf. 1030.53 kW in Table 9) from 151°C to the operating temperature of the methanol synthesis reactor, 250°C.

Table 10 – Reduction of heat demand thanks to thermal integration

	$\dot{Q}_{heating}$ [kW]	$\dot{Q}_{cooling}$ [kW]
<b>W/out Thermal Integration</b>	5645.72	9781.79
<b>With Thermal Integration</b>	1057.30	5193.38

The above-mentioned results are reported using hot and cold composite curves in Figure 6, which highlights the recovery of about 4588 kW corresponding to the range in which the two curves overlap: a design of the network of recovery heat exchangers for the improved energy integration of the plant is reported in Figure S.1 in the Supplementary Material.

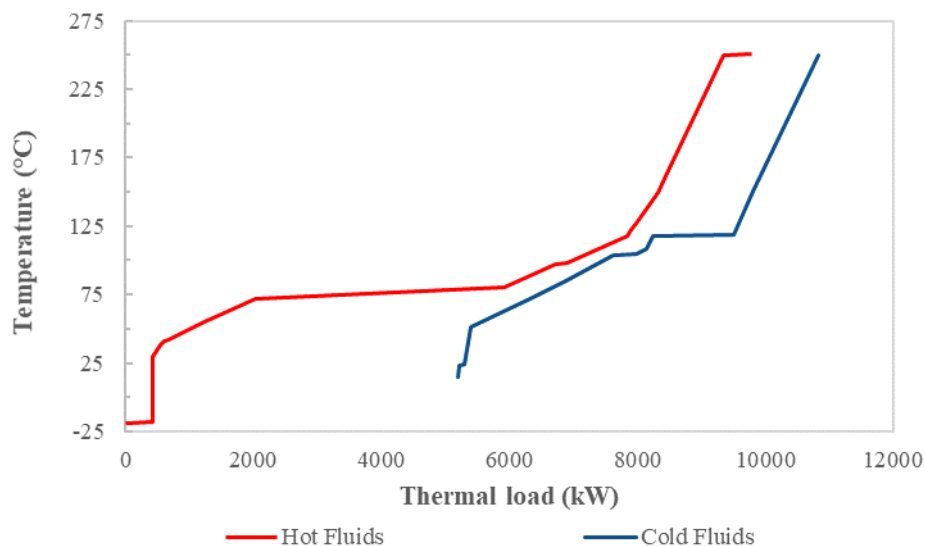


Figure 6 - Hot and cold composite curves resulting from pinch analysis

#### 4.2. Energy and environmental performance

Table 11 reports the values of the KPIs defined in Section 3.4 to assess the performance of the Power-to-MeOH system from energy and environmental standpoints. The capture of a constant flow of about 1.25 t/h of CO<sub>2</sub> from the coal-fired plant exhausts and the exploitation of renewable energy to run the plant ensure an annual emission reduction to the atmosphere of 9968 tons of CO<sub>2</sub>.

The annual water consumption resulting from the simulations run is about 12,635 m<sup>3</sup>, of which:

- 1005 m<sup>3</sup> (ca. 8%) to refill water in the carbon capture solvent due to evaporation and entrainment;
- 8593 m<sup>3</sup> (ca. 68%) to feed electrolysis;
- 3037 m<sup>3</sup> (ca. 34%) as fluid fulfilling the cooling demand of the plant.

The estimation of the different shares highlights that water electrolysis is the most consuming section not only for what concerns electric energy, but also water. Following a Power-to-MeOH scheme through the electrochemical conversion of renewable energy into hydrogen, this issue is unavoidable and must be carefully accounted for when assessing the plant location and the consequent impacts on water availability in different socio-economic contexts.

The benefits of the thermal and mass integration become evident also when analyzing the SPECCA. The specific energy consumption of carbon capture section without thermal integration (i.e. considering power consumption and heating demand associated to the components of carbon capture section in Table 9) is around 1.25 kWh/kgCO<sub>2</sub>. This value is in line with the typical values of traditional post-combustion applications in coal-fired power plant,

0.72-1.53 kWh/kg<sub>CO2</sub> [33] for the regeneration of a commercial 7m MEA solvent. The integration of the sections reduced the thermal demand of the full Power-to-MeOH chain, avoiding the need of an external heat source for carbon capture section thanks to the network of recovery heat exchangers, but limiting thermal supply to the heater at the inlet of methanol synthesis reactor (H-301). In this way,  $Q_{th_{CO_2}}$  in (Eq. 17), accounting mainly for the regeneration heat duty at the stripper reboiler, becomes 0 and the SPECCA drastically reduces to 0.094 kWh/kg<sub>CO2</sub>.

The global efficiency of the plant accounts for the transformations occurring in all the sections of the Power-to-MeOH conversion process. The reduction from the 42.48% gross efficiency reported earlier to 37.22% (Table 11) is mostly due to the residual heating demand after thermal integration, with a smaller influence of power consumption of compressors and pumps. It is worth remarking that  $\eta_{gl}$  would be equal to 26.74% in lack of thermal integration: the 81% reduction of heating demand thanks to thermal integration allows a strong improvement of the system performance.

Table 11 – Key Performance Indicators

Key Performance Indicator	Unit	Value
Reduction of carbon emissions to the atmosphere, $m_{CO_2}$	t/y	9968
Water consumption, $V_{H_2O}$	m <sup>3</sup> /y	12.63*10 <sup>3</sup>
SPECCA (w/out thermal integration)	kWh/kg <sub>CO2</sub>	1.248
SPECCA (with thermal integration)	kWh/kg <sub>CO2</sub>	0.094
Global efficiency of the Power-to-MeOH plant, $\eta_{gl}$	-	37.22%

#### 4.3. Results of the techno-economic assessment

Starting from the results of the simulation of the steady-state operation of the plant, we carried out a techno-economic analysis following a bottom-up approach for the determination of the Total Overnight Cost under three different techno-economic scenarios (1-optimistic, 2-realistic, 3-pessimistic) and the operating costs considering different RES to fulfil the plant electricity demand. The following sections report these results and the main indicators of plant economic viability identified in Section 3.4: COM (cost of methanol) and IRR (Internal Rate of Return) with its levelized cost of methanol (LCOM), investigating the possibility of selling electrolytic oxygen.

#### *4.3.1. Estimation of the Total Overnight Cost*

The size of each component was determined according to the results of the simulations run with Aspen Plus™ at the nominal operating point of the plant. Following the bottom-up procedure described in Section 3.3.1, we estimated the bare module cost of each of the major components in the plant (represented in the scheme in Figure 1) and we defined a confidence range of the TOC of the plant introducing the three economic scenarios. The TOC of the 788-kg/h power-to-MeOH plant is between 27.7 M€ (scenario 1-optimistic) and 40.2 M€ (scenario 3-pessimistic), with a realistic estimate of 31.1 M€, of which we can allocate:

- 6% to carbon capture section;
- 43-45% to water electrolysis section;
- 5-6% to methanol synthesis section;
- 43-45% to the TOC of the network of recovery heat exchangers (including heaters and coolers) resulting from the pinch analysis and designed in Figure S.1 in the Supplementary Material.

Although increasing the initial investment cost, the realization of the heat recovery network allows to reduce the operating costs for the correct thermal management of the plant, exploiting the available resources in a smarter way. The fulfilment of heating and cooling demands with external heat sinks and sources would increase the OPEX.

The cost of a scheme without any thermal integration (like in Figure 1) would probably be lower and allocated in the different sections. Also, in this case, the main part of the plant TOC would be allocated to the water electrolysis section due to the high specific cost of the electrolyzer. Even though alkaline stacks are the most mature technology for water electrolysis, the volumes installed worldwide for stationary conversion and/or storage applications are still limited. Incentives for the commercial use of electrolytic hydrogen as a vector for the long-term storage of RES would be of vital help for a rapid cost decrease, hence making systems like the examined Power-to-MeOH plant more economically viable.

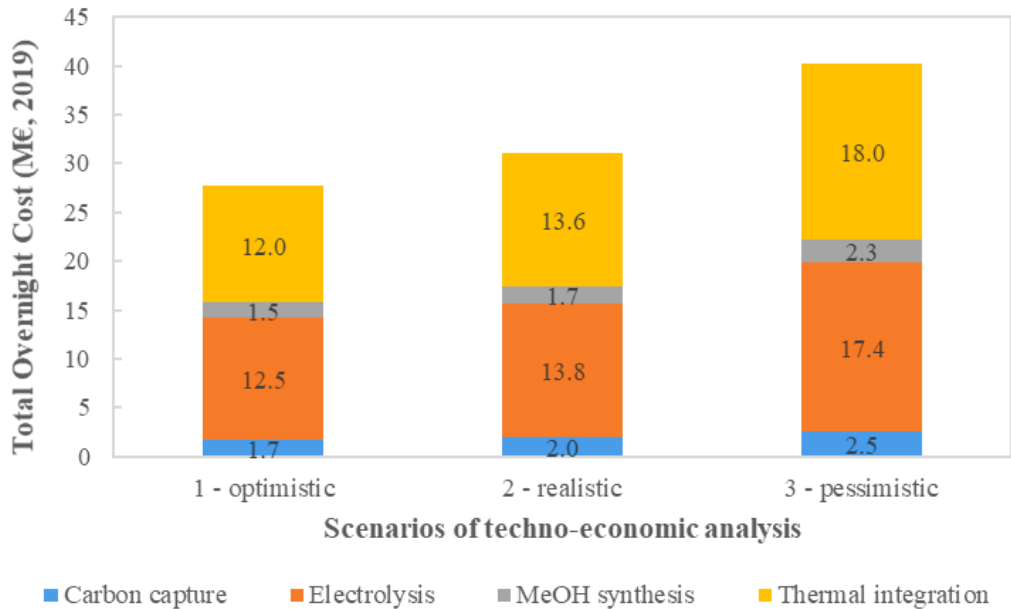


Figure 7 - Contribution of each section to the CAPEX in the different techno-economic scenarios

#### 4.3.2. Estimation of the Operating Costs

The Operating Costs (OPEX) of the whole plant contributing to the definition of the Total Investment Cost (TIC, Eq. 16) in the cash flow analysis were divided in fixed and variable costs. In particular, the fixed costs include the maintenance and the insurance costs; variable costs include costs  $OC_{1-7}$  reported in the lists in Section 3.3.2. The estimation of the maintenance and the insurance costs as percentage of the Total Plant Cost (TPC) justifies the variation of the fixed OPEX in the three techno-economic scenarios, as reported in Table 12. The item “Variable OPEX” in the same table includes:

- 32.7 k€/year for water supply to the different processes ( $OC_2$ );
- 10.2 k€/year for cooling water ( $OC_3$ );
- 4.8 k€/year for MEA refill ( $OC_5$ );
- 130.8 k€/year for the electrolysis stack replacement ( $OC_6$ , supposing to amortize the expenses at 11<sup>th</sup> and 21<sup>st</sup> year over the whole plant lifetime of 25 years);
- 3 k€/year for the replacement of the methanol synthesis catalyst ( $OC_7$ ).

The last operating costs  $OC_1$  and  $OC_4$  are associated to the consumption of electric energy. As explained earlier, we investigated different scenarios in which the required power supply comes from different renewable energy

sources (cases A-G listed in Table 8), each with its average cost. Hence, the cost of feeding the 94.13 GWh/year to run the plant ranges from a minimum value of 3.86 M€/year of hydropower (case C) to a maximum value of 15.2 M€/year in case of CSP (concentrated solar power, case E). This range reflects the different levels of technology readiness, maturity and availability of the two solutions. In any case, feeding the power demand of the plant represents the main operating cost: thus, the choice of the source to produce renewable energy has a strong impact of the economic viability. The cheapest of the new-generation RES is onshore wind (case G): its cost is just 19% higher than hydropower. While hydropower can be installed only in locations with suitable water basins and watercourses ensuring a certain level of availability, wind is a resource that is more distributed worldwide. However, a suitable hydrogen buffer charging and discharging continuously would be required to face the power demand-generation mismatch and the unpredictable variations of wind, hence with an effect on the costs: this is out of the scope of this work and we assume that renewable energy surplus is always available there to fulfil plant power demand, without modeling it.

For each of the A-G cases, we can define an estimate of the OPEX in the three techno-economic scenarios. These values range from 5.12 M€/year of the scenario C.1 (hydropower with optimistic techno-economic hypotheses, i.e., scenario 1) to the upper boundary of 16.8 M€/year of the scenario E.3 (where the system power demand is fulfilled by a CSP plant, under techno-economic assumptions of scenario 3-pessimistic).

Table 12 – Operating costs of the Power-to-MeOH plant

Costs (k€/year)		Techno-economic scenario		
		1 – Optimistic	2 – Realistic	3 – Pessimistic
Fixed OPEX		938	1034	1306
Variable OPEX (OC <sub>2+3+5+6+7</sub> )			181.5	
Scenario for electric energy supply (OC <sub>1+4</sub> )	A		5097	
	B		5919	
	C		3864	
	D		6988	
	E		15208	
	F		10441	
	G		4604	

### 4.3.3. Cost of methanol

Once determined the TIC with the estimates of TOC and OPEX, we computed the cost of methanol (COM), defined as the production cost of methanol that ensures a breakeven over the project lifetime of 25 years (Eq. 19). Considering each source of renewable energy coupled with the plant (cases A-G), we defined a range of COM according to the techno-economic hypotheses presented earlier (scenarios 1-3).

The results are reported in Figure 8, where blue bars represent the range of variation between pessimistic and optimistic scenarios. The values obtained under the realistic techno-economic assumptions (scenario 2) are explicitly reported for the sake of clarity and marked with a blue dot. It is important to remark that this estimation still does not account for the sale of electrolytic oxygen, which is instead vented. We compared the values of COM with the current and the future market price of methanol, 400 €/t and 800 €/t respectively [62].

As expected from the results of Section 4.3.2, case C (hydropower) presents the lowest COM, ranging from 823 to 906 €/t. This range is very close to the predicted price of methanol on future market, hence making this system a competitive configuration for the decarbonization of methanol-based industry and market even without any valorization of electrolytic oxygen.

Five of the considered RES gather in a range that is not far from 800 €/t: in order of increasing price we find hydropower, onshore wind (941-1023 €/t), bioenergy (1019-1101 €/t), geothermal (1150-1232 €/t) and solar PV (1319-1401 €/t). Cases G (onshore wind) and D (solar PV) are of particular interest because they would allow the storage and conversion of large amounts of non-programmable energy generation, hence offering a solution to the abovementioned issues of RES mismatch or, even worse, avoiding its curtailment.

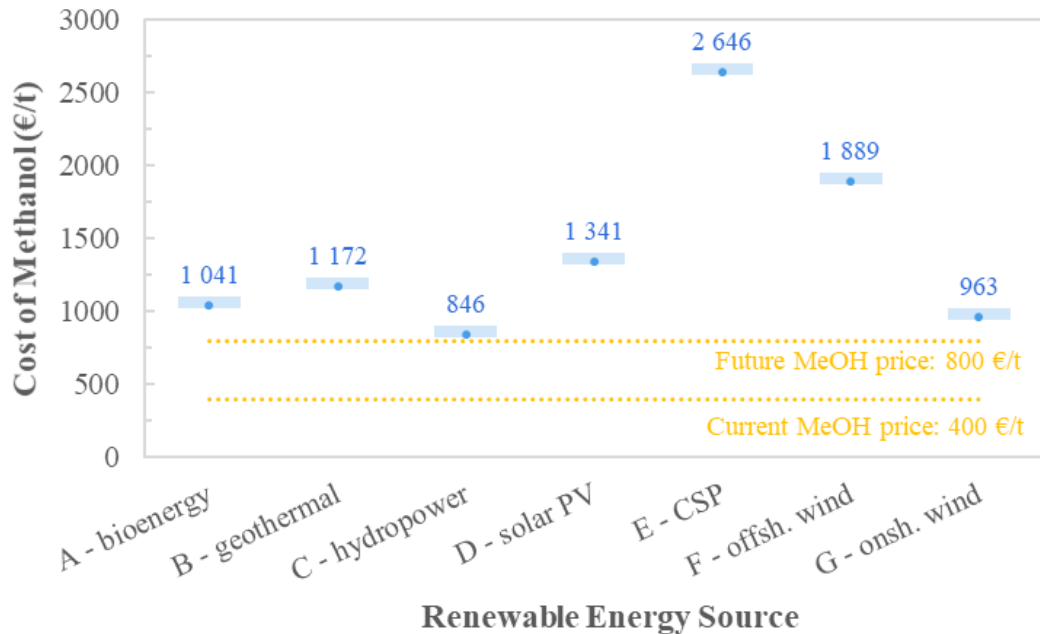


Figure 8 - Variation of COM with the techno-economic scenario and the source of renewable energy

The results obtained strictly depend on the economic assumptions considered. More than others, the cost of electric energy used to run the plant: reductions of this input parameter in the range reported in Table 8 [59] would draw the COM estimates closer to the current methanol price of 400 €/t. For instance, COM would fall under the 1000 €/t threshold whether the lower boundaries of electric energy cost in Table 8 were valid for hydropower (601-684 €/t), onshore wind (785-866 €/t), bioenergy (837-919 €/t), solar PV (967-1049 €/t) geothermal (993-1075 €/t). Case C-hydropower would entirely fall in the 400-800 €/t band, confirming its economic viability under current economic conditions. Finally, a detailed forecasting of the average cost of electric energy would establish the economic competitiveness of these systems in the future methanol market.

The two remaining cases rely on technologies that are still not diffused for commercial application, hence resulting in high values of COM: offshore wind (1867-1949 €/t) and concentrated solar power (2624-2706 €/t). Further demonstration and larger installation would decrease the average cost of electricity from these sources, hence reducing the gap with the cheaper technologies. It is worth remarking that we considered global weighted-average costs in order to be as general as possible without any preference for plant location: local variations of these costs would push up and down the COM confidence bars. However, they represent the least favorable options for this Power-to-MeOH under the current economic conditions. Even considering the lower boundaries of electric energy costs in Table 8, they are too expensive to get a COM smaller than 1600 €/t (i.e. twice the future methanol price of 800 €/t).

#### *4.3.4. Influence of the sale of oxygen from water electrolysis*

So far, we have considered methanol as the only product that have a market value. However, oxygen is produced in a great amount as a by-product of water electrolysis (ca. 8 kg per kg of hydrogen). When compressed or liquefied and stored in a tube trailer (with purchasing and operating costs that are neglected in our techno-economic assessment), oxygen can be used for many industrial and manufacturing applications, thus representing a second income in the cash flow analysis.

This section presents our investigation about the effect of the option of selling oxygen on the economic performance of the plant, considering three costs (0-70-150 €/t) in order to span a range for the economic KPIs with enough confidence. The cost of 0 €/t corresponds to the base scenario of oxygen venting.

The income from the sale of oxygen partially counterbalances the OPEX of the plant. In other words, part of the revenues entirely charged on the sale of methanol in Section 4.3.3 can be now allocated to the optional sale of oxygen. The main effect is the reduction of the COM, with the same annual production of methanol (Eq. 19), as shown in Figure 9. For the sake of simplicity, we represented the effect only for the most and the least favorable RES scenarios resulting from Section 4.3.3 (C-hydropower and E-concentrated solar power), with the other configurations being included between these boundaries.

With respect to the values of 824-906 €/t reported in Section 4.3.3 for case C, the COM reduces by 14% (703-784 €/t) and 30% (565-647 €/t) when oxygen is sold at 70 €/t and 150 €/t, respectively. The option of selling oxygen has the beneficial effect of pushing the COM confidence range down in the 400-800 €/t band, between the current and the future market price of methanol. Case C is not only the most viable configuration for the generation of renewable methanol, but would be also competitive with the current “fossil-based” technology chains characterized by an average price that is susceptible to continuous market fluctuations and has overcome the threshold of 400 €/t many times in US and European markets [63] in recent years.

If the cost of electricity in case C was 26.2 €/MWh (lower bound in Table 8), the COM would furtherly reduce to 481-563 €/t (O<sub>2</sub>: 70 €/t) and 343-425 €/t (O<sub>2</sub>: 150 €/t): in this optimistic – but not unreal – scenario, the cost of the methanol produced by the examined system would be even lower than the average value considered for current market. Similar trends and qualitative results can be expected for the other scenarios of the low-COM block (onshore wind , bioenergy , geothermal , solar PV) identified in Section 4.3.3.

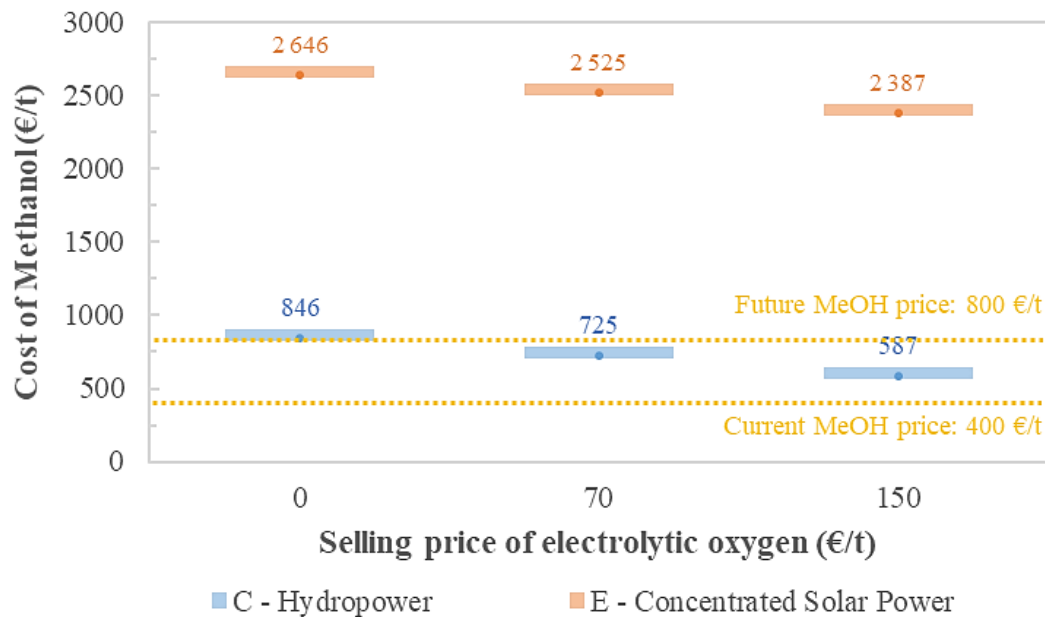


Figure 9 - Effect of the sale of oxygen on the production cost of methanol (COM)

The COM estimates for the cases F-offshore wind and E-concentrated solar power (reported in Figure 9 as red bars) are extremely high. The weight of high-cost electric consumption on the OPEX limits the effect of oxygen sale on the COM reduction with respect to the 2624-2706 €/t in case of venting: -4.5% (O<sub>2</sub>: 70 €/t) and -10% (O<sub>2</sub>: 150 €/t). Even with the option of oxygen sale as a second income, these configurations keep being out of the methanol market.

#### 4.3.5. Internal Rate of Return: a sensitivity analysis

We finally provide a sensitivity of the main indicator of the investment profitability (i.e., the internal rate of return, IRR) to the techno-economic scenarios (1-optimistic, 2-realistic and 3-pessimistic). In particular, we assessed the levelized cost of methanol (LCOM) ensuring an internal rate of return in the range 0-10% in the two extreme RES configurations (cases C-hydropower and E-concentrated solar power with the respective average values of LCOE in Table 8), considering the option of selling oxygen at different prices: 150 €/t in scenario 1, 70 €/t in scenario 2, 0 €/t in scenario 3.

Figure 10 highlights the results of the sensitivity analysis, with the LCOM associated to IRR = 0% equal to the COM estimated in different scenarios (Figure 9), by definition. As expected, the higher the LCOM, the more profitable the investment is and, hence, the higher the value of the IRR is. What is significant is the comparability

of the LCOM with the future market price of methanol even at very high values of IRR in case C. The lower bound of the blue confidence range in the following figure (corresponding to LCOM in scenario C.1 with oxygen sold at 150 €/t) exceeds 800 €/t at IRR = 8%, a value associated to a very good return performance of the investment. The values of LCOM in realistic scenario (C.2 with oxygen sold at 70 €/t) keep under the threshold of 1000 €/t, except for the value associated to IRR = 10% (1072 €/t).

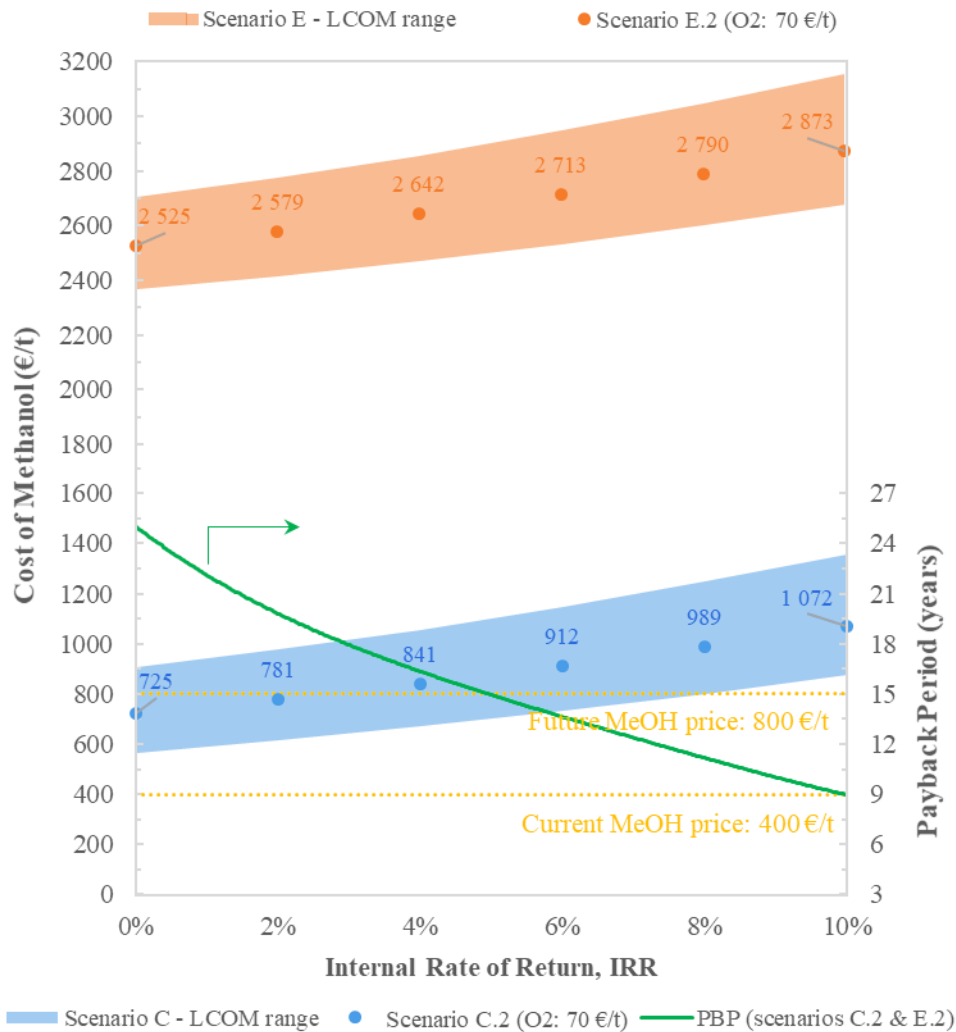


Figure 10 - Variation of the Levelized Cost Of Methanol (left) and Payback Period (right) with the investment internal rate of return in the two extreme scenarios (C-hydropower and E-concentrated solar power)

However, if the cost of electric energy from hydropower was 26.2 €/MWh, the investment IRR would be 10% with

- LCOM = 655 €/t for scenario C.1 with oxygen sold at 150 €/t;
- LCOM = 853 €/t for scenario C.2 with oxygen sold at 70 €/t;
- LCOM = 1135 €/t for scenario C.3 with oxygen vented.

With such IRR, the investment would have an excellent return performance, being extremely competitive in scenarios C.1 and C.2 and with the possibility of a good benefit-cost ratio: if the average discount rate valid during project lifetime was smaller than the IRR, the net present value (NPV) would be strictly positive with certain earnings at the end of plant life.

The results of the sensitivity analysis justify again the option of selling oxygen: its sale determines an average reduction of LCOM of 20% from scenario C.3 (with oxygen venting) to scenario C.1 (O<sub>2</sub>: 150 €/t), hence pushing the LCOM confidence range down in the 400-800 €/t band at lower values of IRR.

As expected from the results of LCOM, configuration E is even less competitive if a higher investment IRR is aimed: the prohibitive cost of electric energy would require an excessively high LCOM (2675-2873 €/t at IRR = 10%).

We finally provide the variation of the Payback Period (PBP, Eq. 21) in the different scenarios: green curve (right-hand axis in Figure 10) is provided for scenarios C.2 and E.2 (both with O<sub>2</sub> sold at 70 €/t), but the trend is almost the same for all the other cases. Since the PBP does not account for the discount rate (i.e. the time value of money), the configurations with COM ensuring an IRR = 0% are characterized by a PBP of 25 years, i.e., the project lifetime. Selling methanol at higher COM ensures higher values of IRR and, hence, smaller payback periods: a minimum PBP of 9 years is reached for configurations with IRR = 10%. The graph also provides a qualitative information about the discounted payback time (PBT): if the discount rate was, for instance, 4%, the LCOM should be higher than 841 €/t in scenario C.2 to have a PBT smaller than plant lifetime and a positive NPV. Otherwise, the investment would not return.

Hence, the results of the sensitivity analysis performed can be interpreted as a profitability map for the investment in the Power-to-MeOH plant: for each techno-economic and energy scenario, we provided the indicators of the investment profitability (IRR) and a comparison of the corresponding LCOM with the predicted cost range of the future methanol market, hence outlining the conditions for the system to be competitive.

## 5. Conclusions

We assessed an integrated system converting about 1.25 t/h of CO<sub>2</sub> captured from a coal-fired power plant into 788 kg/h of methanol through the reaction with “green” hydrogen from water electrolysis (10 MW), following the Power-to-MeOH route. A comprehensive model to simulate the operation of the different sections (water

electrolysis; carbon capture; methanol synthesis) was developed to determine the mass and energy balances of the plant and assess its energy and environmental performance.

- The system ensures an annual reduction of about 10,000 tons of CO<sub>2</sub> with a water consumption of 12.63 \* 10<sup>3</sup> m<sup>3</sup>/y.
- Heat recovery through the thermal integration of the sections (network of heat exchangers designed in Figure S.1 of the Supplementary Material) can heavily reduce the heating and cooling demand of the plant by 81% and 47%, respectively from the initial values of 5.65 MW and 9.78 MW.
- The specific energy consumption for CO<sub>2</sub> avoided (SPECCA) of 1.26 kWh/kg<sub>CO2</sub> for the system without thermal integration perfectly agrees with the literature values. Heat recovery through the different sections of the plant allows to reduce dramatically the thermal penalty for the solvent regeneration, hence leaving a residual energy consumption of 0.094 kWh/kg<sub>CO2</sub> for the carbon capture section.
- The 81% reduction of heating demand thanks to thermal integration allows a strong increase of the system global efficiency from 26.74% to 37.22%, much closer to the value of the Power-to-MeOH gross efficiency (42.48%).

The mass and energy balances resulting from the model simulations were the input for a detailed techno-economic assessment based on a bottom-up approach. From the estimated size of each component, we computed the plant Total Overnight Cost (TOC) under the hypotheses of three economic scenarios: 1-optimistic, 2-realistic, 3-pessimistic. To complete the cash flow analysis, we defined and computed a list of operating costs (OPEX): we assessed their sensitivity to the cost of input electricity introducing seven renewable energy sources (RES) to feed electrolysis and the Balance of Plant (scenarios A-G).

- We estimated a TOC between 27.7 M€ (scenario 1-optimistic) and 40.2 M€ (scenario 3-pessimistic), of which 85-90% can be allocated to the realization of water electrolysis section and the network of recovery heat exchangers for thermal integration.
- The fixed maintenance and insurance costs range from 0.9 to 1.3 M€/y. Variable OPEX are equal to 181.5 k€/y, to which one should sum the cost of feeding the 94.13 GWh/year to run the plant: this contribution ranges from the minimum of 3.86 M€/year in case of hydropower to an upper limit of 15.2 M€/year in case of CSP (concentrated solar power).
- Considering each source of renewable energy coupled with the plant, we defined a confidence range of the cost of methanol (COM) according to the techno-economic hypotheses of scenarios 1-3. Five of the considered RES gather in a range with a COM not far from the future market price of methanol (800 €/t), which are in order of increasing cost: hydropower (823-906 €/t), onshore wind (941-1023 €/t), bioenergy (1019-1101 €/t), geothermal (1150-1232 €/t) and solar PV (1319-1401 €/t). The two remaining scenarios

are characterized by high values of COM: offshore wind (1867-1949 €/t) and concentrated solar power (2624-2706 €/t).

- We investigated the effect of the option of selling oxygen on the economic performance of the plant. With respect to the previous values of 823-906 €/t, the COM in case of hydropower reduces by 14% (703-784 €/t) and 30% (565-647 €/t) when oxygen is sold at 70 €/t and 150 €/t, respectively: selling oxygen has the beneficial effect of pushing the COM confidence range closer to current MeOH market price (400 €/t), improving the competitiveness of the examined system with the traditional methanol synthesis technology chains. Contrarywise, the COM reduction with respect to the 2624-2706 €/t when oxygen venting is limited to -4.5% (O<sub>2</sub>: 70 €/t) and -10% (O<sub>2</sub>: 150 €/t) in case of concentrated solar power.
- We finally provided a map of the profitability of the investment in the examined Power-to-MeOH plant: we estimated the levelized cost of methanol (LCOM) ensuring an internal rate of return ranging from 0 to 10% for each of the techno-economic scenarios identified. Assuming 10% as the desired IRR, LCOM spans from 874 to 1356 €/t in case of hydropower, hence close to the future market price of MeOH with margin of improvement (655-1135 €/t) in case of lower costs of electric energy from hydropower (26.2 €/MWh). Hence, we proved the techno-economic feasibility of employing “green” methanol as a chemical vector for the decarbonization of process industry even in realistic economic scenarios.

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