POLITECNICO DI TORINO Repository ISTITUZIONALE

Techno-economic and exergy analysis of polygeneration plant for power and DME production with the integration of chemical looping CO2/H2O splitting

Original

Techno-economic and exergy analysis of polygeneration plant for power and DME production with the integration of chemical looping CO2/H2O splitting / Uddin, Azhar; Ferrero, Domenico; Santarelli, Massimo. - In: ENERGY CONVERSION AND MANAGEMENT. - ISSN 1879-2227. - ELETTRONICO. - 186:(2019), pp. 200-219. [10.1016/j.enconman.2019.02.043]

Availability: This version is available at: 11583/2726896 since: 2021-02-14T09:36:04Z

Publisher: Elsevier

Published DOI:10.1016/j.enconman.2019.02.043

Terms of use:

This article is made available under terms and conditions as specified in the corresponding bibliographic description in the repository

Publisher copyright Elsevier postprint/Author's Accepted Manuscript

© 2019. This manuscript version is made available under the CC-BY-NC-ND 4.0 license http://creativecommons.org/licenses/by-nc-nd/4.0/.The final authenticated version is available online at: http://dx.doi.org/10.1016/j.enconman.2019.02.043

(Article begins on next page)

1

2	Techno-economic and exergy analysis of polygeneration plant for power and DME production
3	with the integration of chemical looping CO_2/H_2O splitting
4 5	Azharuddin Farooqui ^{*¥} °, Felice Di Tomaso*, Archishman Bose ^{\$} , Domenico Ferrero*, Jordi Llorca [¥] , Massimo Santarelli*
6 7	* Energy Department (DENERG), Politecnico di Torino, Corso Duca Degli Abruzzi 24, Torino 10129, Italy
8 9 10	[¥] Institute of Energy Technologies, Department of Chemical Engineering and Barcelona Research Center in Multiscale Science and Engineering, Universitat Politècnica de Catalunya, EEBE, Eduard Maristany 10-14, Barcelona 08019, Spain
11 12 13	^{\$} Marine and Renewable Energy Research (MaREI) Centre, Environmental Research Institute, School of Engineering and Food Science, University College Cork, Lee Road, Cork, T23 XE10, Ireland
14	° Corresponding author.

15 Abstract

In this paper, we present a novel polygeneration plant with carbon capture for the combined 16 power and dimethyl ether (DME) production. The plant layout integrates a chemical looping 17 CO_2/H_2O splitting (CL) unit producing syngas (CO and H₂) for the DME synthesis using the 18 19 exhaust gases of an oxyfuel power cycle. The primary power is generated by oxy-combustion of syngas generated by the reduction of the metal oxide in the reduction unit of the CL redox cycle 20 with incoming natural gas. The oxyfuel power plant also generates steam for combined power 21 production with two streams Rankine cycles. The aim of the present work is to assess the process on 22 the basis of energy and exergetic efficiency and economic performance of the integrated CL unit for 23 combined power and DME production. The integration proposed resulted in a production of 103 24 MW_e and 185.6 ton/day (2.15 kg/s) of DME. The corresponding energy and exergetic efficiency 25 was 50.2% and 45%, respectively. A discounted cash flow analysis was performed to evaluate the 26 profitability of the process. With a carbon credit of \$80/tonne, the plant would be able to meet the 27 current electricity with carbon capture and DME prices of \$50/MWh and \$18/GJ respectively. The 28 economic analysis provided information on the main economic drivers associated with the high 29 capital investment in the process plant with individual sub-systems. The analysis highlighted the 30 31 strong potential of integrating chemical looping CO₂/H₂O splitting for syngas production into polygeneration systems to increase the overall efficiency while reducing the cost of carbon capture. 32

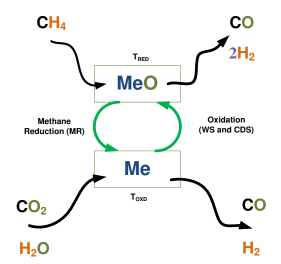
Keywords: CO₂/H₂O splitting, Chemical looping, Polygeneration, Oxyfuel combustion, Dimethyl ether (DME), carbon capture.

35 1. Introduction

CO₂ emissions from the energy sector – mainly from the combustion of fossil fuels – comprise 36 the largest fraction of the global anthropogenic GHG emissions, representing 58% of the total 37 emissions, as of 2014 [1]. The quest to meet the never-ending energy demand and the rise of 38 emissions is leading to the search for innovative technologies and non-petroleum based alternative 39 fuels which would help in restricting the global warming to 1.5°C above the pre-industrial 40 temperatures (new target set by the recent report by Intergovernmental Panel on Climate Change 41 42 (IPCC), 2018) [2]. Among the multiple pathways proposed for the reduction of anthropogenic emissions of CO₂, Carbon capture and utilization (CCU) to convert captured CO₂ into valuable 43 products have recently gained much focus as an alternative to Carbon Capture and Storage (CCS) 44 [3-5]. CCU is not only complementary to CCS in some respects but also provides multi-product 45 outputs through the recycling and reuse of the captured CO_2 in several synthesis processes [6–9]. 46 Carbon capture in power plants comes with huge energy penalty and loss of efficiency. For 47 example, in oxyfuel natural gas combined cycle (NGCC) power plants, Air separation unit (ASU), 48 used to produce pure oxygen for combustion results in the decrease the efficiency of the 49 conventional NGCC by as much as ~13% [10]. Multiple studies have been proposed to gain this 50 loss of efficiency by alternate methods like the use of ion transport membranes (ITM) for oxygen 51 separation instead of ASU [11] or recycling exhaust flue gas to run a redox cycle with a metal oxide 52 [12]. Polygeneration systems, which can combine efficiently multiple utility outputs (e.g., electrical 53 54 power, chemicals, fuels etc.) from one or more input in a single system, provide an interesting option for CCU [13]. Besides the potential to gain significant efficiency and local use of the 55 captured CO₂, suitable integration and synergy between different processes also ensure higher 56 flexibility of operation. This would allow varying between the share of products according to their 57 value, for example, related to fluctuating market prices [14]. Multiple configurations of 58 59 polygeneration systems integrated with CO₂ capture processes have been reported in the literature. Li et al. [15] modelled a polygeneration plant with CO₂ capture for production of power and 60 synthetic natural gas, the proposed arrangement achieving a lower life-cycle energy use and GHG 61 emission with respect to the ultra-supercritical coal power plant. Bose et al. [16] studied a cost-62 63 effective production of urea and power combined with CCS using coal gasification. Jana et al [17] reported the improved sustainability through life cycle assessment for a rice-straw based power, 64 ethanol, heating and cooling polygeneration power plant. Huang et al [18] highlighted both 65 energetic and economic benefit of a coal based polygeneration system for power and methanol 66 production as opposed to single coal-to-methanol or coal-to-power systems. Salkuyeh [19] proposed 67 a novel methanol, DME and power production plant from the combined use of coal and natural gas 68 via chemical looping combustion that not allowed complete carbon captured at improved efficiency. 69

Most polygeneration systems designed or proposed till date have employed coal as the fuel [13]. However, oxyfuel combustion using gaseous fuels like natural gas and biomethane has been shown to be the most promising among the low emission technologies (LETs) [12]. Above such, innovative methods for the use of natural gas are being proposed to improve upon the efficiency of natural gas combined cycle power plants, which can reach an efficiency of as high as 57% [20]. One such innovative technology is the chemical looping CO_2/H_2O splitting cycle using methane reduction, to produce CO and H₂ [12]. This indeed forms an interesting alternative to the solar

77 thermochemical redox cycle, which has gained attention for CO₂/H₂O splitting to produce syngas (CO/H₂) utilizing the successful demonstration of water splitting by oxygen careers [21–24]. For the 78 solar-thermochemical redox cycle, the metal oxide (also called as oxygen career (OC)) undergoes 79 endothermic thermal reduction (TR) step, thus requiring a very concentrated heat, usually provided 80 by solar energy under a very high vacuum pressure [25,26]. In the second step, the reduced oxygen 81 carrier undergoes oxidation by the incoming CO₂/H₂O producing CO/H₂, thereby ensuring a 82 83 continuous cycle. The second step is exothermic in nature and operates at near atmospheric pressures and lower temperature than the thermal reduction creating a temperature and pressure 84 swing between the two steps. Fuel reduction with methane, however, results in the reduction step to 85 operate at near atmospheric conditions, thus, avoiding pressure swing operation. Furthermore, it 86 lowers the reduction temperature, preventing a large temperature swing between the two-steps. An 87 added advantage is gained from the production of syngas in both the steps of redox cycle 88 [12,27,28]. Chemical looping (CL) cycle driven by methane reduction (which can be replaced by 89 bio-methane in future), has an added benefit of being able to operate round the clock, unlike solar 90 91 driven cycles that are constrained by the fluctuation of the solar energy source. A basic schematic of 92 the methane-driven chemical looping CO_2/H_2O splitting cycle is shown in Figure 1.



93

94

95

Figure 1. A representative schematic of chemical looping syngas production through methane reduction and corresponding splitting of water and carbon dioxide.

96 Multiple metal oxide redox pairs have been studied for the CL application [25,26,29]. Among the materials, cerium (IV) oxide (CeO₂) is considered to be one of the most promising for the 97 present application due to its strong ability to undergo cyclic redox reactions while retaining its 98 99 chemical and structural properties together with high resilience to mechanical stress and agglomeration resistant [30]. Ceria reduction by methane has been investigated by Warren and 100 Scheffe [31]. Results indicate that CeO₂ undergoes complete reduction to Ce₂O₃ above 900°C. 101 Accordingly, the CeO₂/Ce₂O₃ redox pair with reduction of CeO₂ in the presence of methane, and 102 subsequent oxidation with CO_2/H_2O can be described in terms of the following equations (1-3). 103

104 Methane reduction (MR):
$$2\text{CeO}_2 + \text{CH}_4 \rightarrow \text{Ce}_2\text{O}_3 + \text{CO} + 2\text{H}_2$$
 (1)

105 Water-splitting(WS):
$$Ce_2O_3 + H_2O \rightarrow 2CeO_2 + H_2$$
 (2)

106 CO_2 -splitting (CDS): $\operatorname{Ce}_2\operatorname{O}_3 + \operatorname{CO}_2 \to 2\operatorname{CeO}_2 + \operatorname{CO}$ (3)

By optimally combining the ratio of water and CO_2 in the inlet gas mixture to the oxidation reactor and the temperature of reaction, the desired composition of syngas can be obtained, to be subsequently utilized for production of chemicals ((H₂/CO: 1.79) methanol [32,33], (H₂/CO: 2.1) jet fuels [34] and naphtha [34–36], (H₂/CO: 1.76) kerosene and gasoil [36] etc.) through industrial processes.

Dimethyl Ether (DME) is one of the most attractive candidates as a synthetic fuel due to its 112 similarity with diesel. Even though DME has a lower LHV than conventional diesel and its use 113 requires pressurization to maintain it in a liquid state at ambient conditions, its physical properties 114 and chemical structure make it a very interesting fuel. Low NO_x, limited hydrocarbon (HC) and 115 almost no SO_x and particulate emissions during the combustion [37,38], are added advantages of its 116 117 use. However, challenges including the need for re-design of the injector and fuel pump systems, compatibility of sealant materials are current limitations to the complete replacement of diesel by 118 DME in conventional compression ignition engines [37]. In this regard, blending of DME with 119 conventional automotive fuels have been advocated as an important strategy to improve both 120 121 applicability and economic viability of DME in the short term [37,39]. Furthermore, the unique physico-chemical properties of DME would allow its use as a raw material for the synthesis of 122 aromatics, gasoline, olefins and other chemicals besides direct use as an alternative fuel or a 123 substitue to conventional refrigerants [40]. It is to highlight that DME also gained attention in recent 124 times due to its physicochemical properties are similar to liquid petroleum gas (LPG) giving the 125 chance to retrofit the LPG based automotives [41]. However, being a synthetic fuel, the price of 126 DME would be constrained by the cost of feedstock, including its transportation, in the present case, 127 128 natural gas [42]. As per long term future predictions, both diesel and natural gas prices have been projected to rise at an eugal steady rate [43,44]. This, even though would allow the relative 129 economic competitiveness of DME with respect to diesel, would increase its cost nevertheless in 130 future. In this regard, research and development for sourcing methane from biomass as biomethane 131 could potentially improve both environemntal and economic volatility associated with DME 132 synthesis. 133

DME synthesis is generally classified as i) two-step process (indirect) which uses 134 hydrogenation to produce methanol and then dehydration to DME ii) the second method is one-step 135 (direct) process reported to be more efficient which uses bi-functional catalysts. Both the pathways 136 are commercially viable technology and invested by companies such as Haldor Topsoe, Korea Gas 137 Corporation, Air products, JFE Holdings, Toyo, MGC, Lurgi and Udhe [45,46]. Synthesis of DME 138 using syngas (CO and H₂) from CO₂/H₂O splitting can, therefore, present an interesting pathway for 139 the production of clean fuels using an unconventional process [47,48]. Interestingly, the production 140 of DME by the single step process has been shown to significantly improve the overall process 141 efficiency with respect to both methanol production or the indirect DME synthesis process, 142 providing additional economic impetus to its commercial application [49]. 143

Alternative methods to produce syngas by the chemical looping processes has been reported such as chemical looping reforming (CLR), autothermal reforming and chemical looping partial oxidation of methane (CLPOM) [50,51]. CLR and autothermal reforming usually operate at a lower

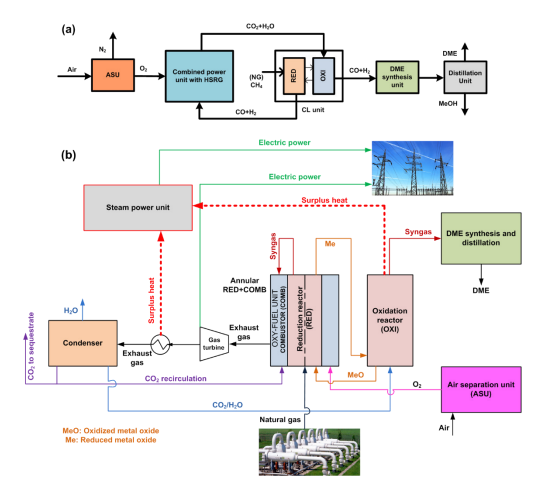
temperature of 800-900°C that produces H₂/CO ratio of 2.8-4.8 and 1.8-4.0 respectively with a 147 higher concentration of CO₂/H₂O at the outlet stream. While syngas production by CLPOM needs a 148 temperature above 1300° C with H₂/CO of 1.7-1.8 and also has lower H₂O/CO₂ in the product [52]. 149 This makes CLR more suitable for Fischer-Tropsch synthesis for methanol or hydrogen production. 150 However, a novel process of generation of syngas from the exhaust stream and re-use within the 151 power plant for producing additional power has been shown to be a viable alternative to improve 152 153 the efficiency with 100% carbon capture [12]. Indeed, within a polygeneration scheme, the use of syngas for synthetic fuel production becomes an imperative option. Hankin and Shah [49] in an 154 study explored the process of DME and methanol synthesis from CO₂ and H₂O. Syngas is produced 155 by water electrolysis and solid oxide electrolysis for CO where all the processes such as DME, 156 methanol synthesis, electrochemical electrolysis and solid oxide electrolysis for CO are investigated 157 by assumption of chemical equilibrium. Salkuyeh and Adam II [19] proposed a polygeneration 158 scheme which combines the coal gasification, natural gas reforming by chemical looping processes 159 such as gasification and combustion to produce power, methanol, and DME. The system was tested 160 161 with iron oxide and nickel oxide oxygen career for chemical looping processes with different gains based on the operability of the system. The path for syngas production as feedstock DME was 162 investigated. 163

164 However, till date, as per the knowledge of the authors, no polygeneration system which integrates the chemical looping CO₂/H₂O splitting (CL) with fuel reduction step and DME and 165 power production have been studied for utility-scale. In this work, an oxyfuel natural gas combined 166 cycle power plant integrated with CL CO₂/H₂O splitting and DME production has been proposed 167 (OXYF-CL-PFG) with a detailed techno-economic, exergetic and environmental assessment. The 168 exergetic study was carried out for the proposed OXYF-CL-PFG layout to identify the sources of 169 irreversibility, with which the proposed layout could be improved and optimized. The analysis 170 includes power production, fuel production, and power consumption, exergy analysis, economic 171 estimation along with the net present value (NPV) with different carbon credit scenarios, as well as 172 efficiency and percentage of carbon captured and recycled. 173

174 **2.** Process and plant description

The proposed polygeneration scheme is an oxyfuel natural gas fed combined cycle power plant 175 integrated with a chemical looping CO₂/H₂O splitting unit (CL) for power and DME production 176 (OXYF-CL-PFG) shown in Figure 2. To maintain the simplicity of analysis, the gas pre-treatment 177 including sulphur removal has been assumed to have occurred upstream [53] The clean natural gas 178 is sent to the chemical looping (CO₂/H₂O) splitting unit where it is converted into a hydrogen-rich 179 syngas by the simultaneous reduction of ceria. The produced syngas is sent to an oxyfuel unit where 180 it is combusted with pure oxygen from an ASU. The hot combustion products, primarily comprising 181 H₂O and CO₂ are firstly expanded in a gas turbine and then sent in a heat recovery steam generation 182 unit (HRSG). Here, the surplus heat is exploited to produce superheated steam for power production 183 in a bottoming steam cycle. Finally, a water condenser partially separates carbon dioxide and water. 184 The large part of the separated CO_2 can be sequestrated for storage or used in other processes, while 185 another fraction together with steam is sent to the chemical looping CO₂/H₂O (CL) unit. In the CL 186 unit, both H₂O and CO₂ are dissociated to H₂ and CO in an oxidation reactor by the reduced ceria 187 from the reduction reactor. The produced syngas from the oxidation reactor is used for DME 188

synthesis. The diluted DME, resulting thus, is cleaned in a distillation unit (or clean-up unit), additionally producing a secondary fuel stream of methanol. In the following sub-sections, the methodology adopted for the presented work, along with more details on each unit and their integration are described.



193

Figure 2: General concept of integration of oxy-fuelled power unit with chemical looping (CO₂/H₂O) splitting unit and DME production process (a) block diagram (b) process flow diagram

196 2.1 Simulation Methodology

The polygeneration plant has been modelled by combining mass and energy balance equations. 197 As per the detailed plant layout proposed in Figure 3. Simulations were performed using the 198 commercial software Aspen Plus v8.8. The characteristic components of the system are the 199 integrated combustor of the oxyfuel unit and the reduction reactor of the CL unit and oxidation 200 reactor, the DME synthesis reactor, the DME distillation columns, and the ASU apart from the 201 standard components of the plant, such as heat exchangers, pumps, compressors, and turbines. The 202 entire modelling was performed with the assumption of chemical equilibrium with the exception of 203 the DME reactor, for which a kinetic approach has been used. Therefore, RGIBBS reactor blocks 204 were used for modelling the oxidation and reduction reactors of the CL unit, as well as the 205 combustor of the oxyfuel unit. The distillation unit and air separation columns were modelled using 206 the RADFRAC column. The DME reactor was simulated with an RPLUG reactor using the 207 Langmuir-Hinshelwood Hougen-Watson (LHHW) kinetic model. During the simulation of this 208

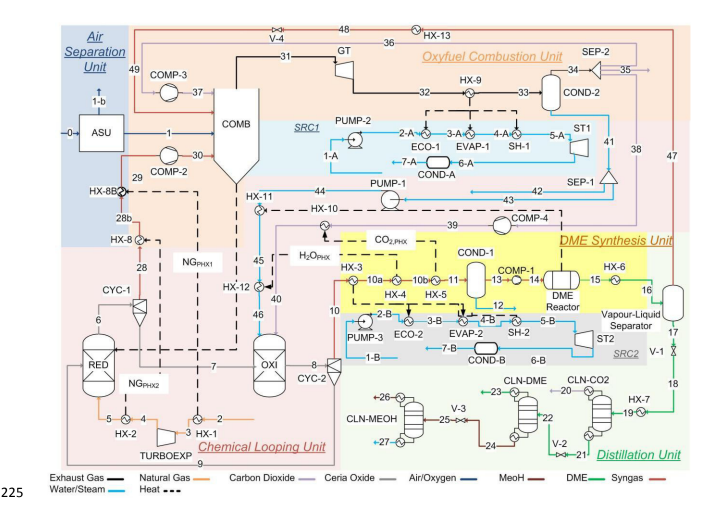
209 component, the Soave-Redlich-Kwong (SRK) EOS model was utilized, which is usually applied to 210 binary components [54]. Graaf et al. [55] demonstrated that the chemical equilibrium of the 211 methanol reaction and water gas shift (WGS) reaction can be well described at high-pressure by 212 using the SRK-EOS model. More details on the modelling approaches followed for the main 213 components of the plant are given in section 2.1-3.

214 The material streams used in the model involve conventional and solid components. The Peng-Robinson-Boston-Mathias (PR-BM) property method was used for conventional components, as 215 this approach was recommended for hydrocarbon processing applications such as gas processing, 216 refinery, and petrochemical processes [56–58]. This method uses the Peng-Robinson cubic equation 217 of state combined with the Boston-Mathias alpha function for all the thermodynamic properties 218 [57]. The oxygen carriers (CeO_2/Ce_2O_3) used for the chemical looping simulation were 219 implemented as conventional pure solid components. For this type of streams, the Barin equation 220 was used [59,60]. The main hypotheses used in the modelling phase are summarized in Table 1. 221 CLN-CO₂, CLN-DME, and CLN-MeOH are the columns used in the distillation unit. 222

Table 1. Main assumptions and hypothesis used in the process simulation.

Natural gas	Composition (std.vol%): 93.1% CH ₄ , 3 LHV=47.1 MJ/kg [61];	2% C ₂ H ₆ , 1.6% N ₂ , 1.1% C ₃ H ₈ , 1.0% CO ₂ ;					
Oxidation and reduction reactors	Model: RGIBBS, no heat losses; 10°C dra assumed in order to assess heat losses; Mod	op for ceria recirculation from OXI to RED was lel: RGIBBS;					
Combustor	Model: RGIBBS; $\Delta P=0.2$ bar, no heat losse	s;					
Compressors, pumps and turbines	$\eta_{is,comp} = 0.9, \ \eta_{mech,comp} = 0.98, \ \eta_{is,pump} = 0.9, \ \eta_{driver,pump} = 0.90, \ \eta_{is,turb} = 0.9, \ \eta_{mech,turb} = 0.98;$						
Oxygen carrier	Solid ceria: CeO ₂ /Ce ₂ O ₃ , diameter=100 μm from OXI to RED;	Solid ceria: CeO_2/Ce_2O_3 , diameter=100 µm; Temperature drop of 20°C during ceria recycling from OXI to RED;					
DME reactor	Model: RPLUG multi-tube reactor, Operati	on: T=250°C P=50 bar;					
Heat exchangers	$\Delta T_{min}=10^{\circ}C;$						
Distillation unit	Model: RADFRAC, Reboiler type: Kettle.						
CLN-CO ₂	CLN-DME	CLN-MeOH					
P=10 bar	P=9 bar	P=2 bar					

224



226

Figure 3. Detailed polygeneration plant layout OXYF-CL-PFG.

227 **2.2 Polygeneration plant units**

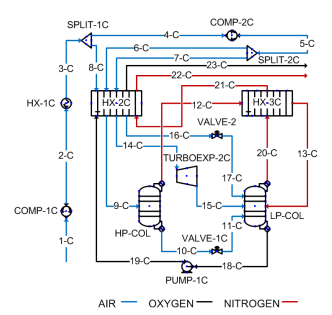
228 2.2.1 Chemical looping CO_2/H_2O splitting unit (CL unit)

The chemical looping unit consists of two interconnected reduction reactor (RED) and the 229 oxidation reactor (OXI) operating at 2 bar with the circulating oxygen carrier pair (CeO_2/Ce_2O_3). 230 The pre-cleaned natural gas, at a grid pressure of 70 bars (stream 2) [62] is heated up at 290°C and 231 expanded to 2 bar via the turbo-expander (TURBOEXP). Table 1 lists the composition of natural 232 gas at the inlet to the plant (without H_2S). The preheating is necessary to prevent an outlet 233 temperature of the natural gas (stream 4) from the turbo-expander lower than 0°C. After the 234 expansion of the natural gas, it is heated to 890°C (stream 5) and fed to the RED. For the 235 endothermic reduction reaction, external heat is mandatory to maintain the reaction temperature. 236 Ceria reduction by methane occurs above 900°C to achieve full conversion to CO and H₂ as well as 237 a reduction to Ce₂O₃ [31]. From the thermodynamic studies, it was found that 40 to 60% excess 238 flow of methane is required to ensure complete conversion of OC below 950°C. It was also found 239 that the most suitable methane to ceria flow ratio (CH₄/CeO₂) for the reduction reactor was 0.7 240 instead of the stoichiometric ratio of 0.5. Here CeO₂ (stream 9), at an inlet temperature of 1312°C 241 as a result of the exothermic oxidation reaction, is completely reduced with natural gas, producing a 242 syngas in a 2:1 H₂/CO ratio (reaction 1) and unreacted natural gas [12]. As for the external heat 243 source to sustain the reaction in the RED, it has been proposed to use a part of the heat generated in 244

the oxyfuel combustion chamber. To this end, a reduction reactor thermally integrated with the 245 oxyfuel combustion chamber was proposed utilizing an annular combustion chamber design already 246 analyzed by Khan and Shamim [63] (see Figure 2 (b)). The hot syngas (stream 6) produced in RED 247 exits it at 900°C and is separated from the solid (stream 7) by a cyclone (CYC-1), cooled and sent 248 to the oxyfuel unit. The reduced ceria is fed, without an intermediate heat recovery, into the OXI 249 where it is then oxidized (reactions 2 and 3) by a gas mixture coming from the oxyfuel unit. The gas 250 251 mixture of 60% H₂O and 40% CO₂ (stream 40 and 46) is necessary to ensure an H₂:CO ratio of 1:1, ideal for DME production, as described in section 3. It is observed that in order to achieve a full 252 oxidation of Ce₂O₃, a 60% excess of the gas mixture is required. Before the oxidation, both water 253 and carbon dioxide is compressed at the operating pressure of OXI (2 bar), respectively with a 254 pump (PUMP-1) and a compressor (COMP-4), and heated up at 500°C. Since the reactions in the 255 oxidation reactor are exothermic and the reactor itself is set as adiabatic, the outlet temperature of 256 the reactor goes to 1322°C. The hot syngas produced is separated from the oxidized ceria by the 257 cyclone separator (CYC-2), cooled down (stream 10, 10a, 10b, 11) and sent to the DME unit, while 258 259 the solid stream is re-circulated back for a new reduction cycle (stream 9).

260 2.2.2 Air separation unit (ASU)

The ASU consists of a cryogenic distillation unit able to produce 99.99% pure O₂. The 261 schematic of the ASU layout is shown in Figure 4. The air is separated into two thermally 262 interconnected distillation columns, HP-COL and LP-COL, which work at 5 and 1.2 bar 263 respectively [64–66]. The overall refrigeration is driven by the expansion from high pressure (30 264 bar) of the compressed air (stream 6-C and 7-C, which become 14-C and 16-C respectively, after 265 cooling down in HX-2C) through the VALVE-2 and the TURBOEXP-2C. The inlet air (1-C) is 266 compressed at 6.3 bars by the compressor COMP-1C and separated in two streams (4-C and 8-C) 267 by the splitter SPLIT-1C. The stream 8-C is cooled down (becoming 9-C in the figure) in the 268 exchanger HX-2C by the cold products (steam 19-C) of the LP-COL and then is sent to the HP-269 COL. The HP-COL is a 40 stages distillation column which produces N₂ -rich gaseous (stream 12-270 C) from the top and an O₂-rich liquid stream (stream 10-C) from the bottom. The latter stream is 271 272 further cooled down through Joule-Thomson effect in the valve VALVE 1-C and fed in the 56 stage low-pressure column. The low-temperature air streams 15-C and 17-C, together with the rich-in O₂ 273 liquid stream 11-C, provide the necessary refrigeration in the LP-COL to obtain pure N₂ (stream 20-274 275 C) from the top and from the bottom condenser the pure O_2 (stream 18-C) is produced. The latter is pumped by the PUMP-1C at the operational condition of the combustor (26 bar) in the oxyfuel unit 276 and heated in the HX-2C to 80°C. The stream data is listed in Table S1 of the supplementary data 277 file. 278



279 280

Figure 4. Detailed layout of the air separation unit

281

282 2.2.3 Oxyfuel combustion unit

In this unit, combustion of syngas is performed with oxygen instead of air. This eliminates the presence of nitrogen in the exhaust gases that would have affected the subsequent CO_2 separation process. Another advantage is a substantial reduction in thermal NO_x due to the absence of nitrogen [67].

287 The primary unit of the oxyfuel unit is the combustor (COMB), where the syngas from the reduction (stream 30) and the non-condensable gases from the clean-up unit (stream 49, mainly CO₂ 288 with CO and H₂) are burnt with a 5% excess oxygen stream derived from the ASU (stream 1). The 289 temperature control within the COMB is maintained by recycling a part of the captured CO₂ 290 (Stream 36). The recirculation ratio was set so as to achieve a combustor outlet gas temperature of 291 292 1377°C in agreement to the limits of the turbine inlet temperature of commercial gas turbines as stated and selected from literature [14,68–70]) while simultaneously providing the required heat 293 sustain the ceria reduction reaction in the RED. The CO_2 and syngas streams entering the COMB 294 are compressed to 26 bar with two two-stage compressors (COMP-2 and COMP-3). The flue gas 295 exiting the combustion chamber is firstly expanded in a two-stage gas turbine GT (26 bar to 5 bar 296 and 5 bar to 1.05 bar) and then sent to the heat recovery steam generator (HX-9) for the generation 297 of steam for the steam power cycle SRC1. More details on the steam power cycle are given in 298 section 2.2.4. Finally, the CO_2 from the exhaust gas is separated from the water in a condenser 299 (COND-2), generating highly pure CO₂ stream that is split into three parts. One part is recirculated 300 301 to the combustor (stream 36), one is sent for sequestration or other applications (stream 35) and the last part (stream 38) is sent to the oxidation reactor for dissociation (OXI). 302

303 2.2.4 Steam power cycle

Two steam Rankine cycles (SRC1 and SRC2) are included in the system layout. The extra heat available within the polygeneration system is exploited to produce steam by heat recovery steam generators (HRSGs), which expands in steam turbines to generate power. The turbines and the

HRSGs were modelled as simple units, without reheating or multi-pressure systems. In fact, as the 307 primary objective of the present study is to understand the benefits deriving from polygeneration by 308 309 integration of a chemical looping unit in a conventional oxyfuel plant, the optimization of the system was not further considered. The SRC1 uses the heat of the flue gases from the oxyfuel unit 310 (stream 32) to produce super-heated steam (125582 kg/h) at 150 bar and 550°C (stream 5A), 311 generating an electrical power of around 44 MW while expanding in the turbine (ST1). The SRC2 312 313 uses the extra heat from the chemical looping unit to produce a smaller flow of steam (8305 kg/h) at the same condition as that of stream 5A (stream 5B), generating 3 MW in ST2. The reason for the 314 choice of two HRSGs connected to two different steam cycles is to ensure flexible operation by 315 minimizing the influence of DME and power production over each other. 316

317 2.2.5 DME synthesis unit

In this unit, the syngas produced in the oxidation reactor (stream 10) is converted into liquid 318 fuel within the catalytic reactor. Before the syngas is fed to the DME reactor, it undergoes 319 condensation (COND-1) to completely remove H₂O at atmospheric pressure. The operating 320 conditions of the DME reactor have been selected from the work of Pozzo et al. [71] fixing the 321 pressure at 50 bar and the temperature at 250 °C. In order to reach the operating pressure of the 322 reactor, the dried syngas (stream 13) is compressed by a three-stage compressor at 50 bar (COMP-323 1). The DME reactor is a fixed bed reactor which is kept at the constant temperature of 250°C by a 324 water-jacket cooler used for saturated steam generation at 2 bar (stream 44) for the oxidation (OXI) 325 reactor. 326

The DME reactor was considered as a multi-tube fixed bed reactor. Each tube contains the dual 327 328 catalyst (physically mixed) with a bed voidage of 0.45. The total density of the catalyst particles is an average of the density of the two catalysts, Cu/ZnO/Al₂O₃, and γ -Al₂O₃, used in the 1:2 optimal 329 ratio. The parameters used for the DME reactor are listed in Table 2. 330

N° tubes	Diamater	Bed	density	density γ-	ρ average	Temperature	Pressure
	[m]	voidage	Cu/ZnO/Al ₂ O ₃	Al_2O_3	$[kg/m^3]$	(°C)	(bar)
			$[kg/m^3]$	$[kg/m^3]$			
5000	0.02	0.45	1200	1470	1380	250	50

332

333 2.2.6 DME distillation unit

The produced DME contains significant impurities, requiring a separation or distillation unit to 334 obtain pure dimethyl ether. The distillation plant comprises a cooling and a gas-liquid separation 335 unit. The cooling unit, represented in the layout by a vapour-liquid separator (VLS), is used to 336 337 produce chilled streams at -40°C resulting in a liquid stream of DME with dissolved CO₂ and MeOH (stream 17) and a gas stream of incondensable gases, namely, H₂, CO, undissolved CO₂ and 338 traces of other diluents (steam 47). The gas stream is re-circulated into the oxyfuel unit and burnt, 339 while the liquid stream is further processed in the gas-liquid separation unit. The gas separation unit 340 is composed of three different distillation columns: CLN-CO2, CLN-DME, and CLN-MEOH (Table 341 3). The first one is used to separate the dissolved CO₂, the second to produce a pure 99% DME and 342 the last one to separate the methanol from the water. Thus, an additional fuel as methanol is 343

344 generated as a by-product of DME distillation. A valve and a heat exchanger are placed before each 345 column in order to adjust the pressure to the optimal value and to have 50% of vapor in the inlet 346 stream [71]. The number of stages used in the distillation columns was estimated by increasing 347 them until a certain change in composition was detected.

Column	T _{REB} [°C]	Q _{REB} [MW]	T_{cond} [°C]	Q _{cond} [MW]	Number of stages	Feed-in stage	Purity of the product [%]
CLN-CO ₂	45.87	1.12	-40.83	-0.64	25	10	-
CLN-DME	150.99	0.93	42.57	-0.55	30	24	99.1
CLN-MeOH	101.53	0.03	66.36	-0.05	24	18	94.1

348 Table 3 Distillation unit operation parameters.

350 *3.* Synthesis of DME

351 *3.1 Reaction scheme*

352 DME production can be realized in two steps (methanol and DME are produced in two different reactors) or in a single step adopting a dual catalyst. The disadvantage of the two-step process is 353 that syngas conversion to methanol is significantly limited by equilibrium and thermodynamic 354 constraints [72]. The conversion of methanol to DME in the single step process shifts the 355 equilibrium toward more methanol production. Consequently, the direct DME synthesis is 356 thermodynamically and economically preferable than the two steps process [73-75] and hence 357 selected in the present study. The overall process can be described by three main reactions: the 358 359 syngas conversion to methanol (reaction (4)), water gas shift (reaction (5)) and methanol dehydration to DME (reaction (6)). 360

$$361 \quad CO_2 + 3H_2 \to CH_3OH + H_2O \quad -\Delta H_{298 \ K:01 \ MPa} = 49.2 \ kJ \ / \ mol \tag{4}$$

$$362 \quad CO + H_2O \to CO_2 + H_2 \quad -\Delta H_{298 \ K:01 \ MPa} = 41.2 \ kJ \ / \ mol \tag{5}$$

$$363 \qquad 2CH_3OH \to CH_3OCH_3 + H_2O - \Delta H_{298 K;0:1 MPa} = 24.0 \ kJ \ / \ mol \tag{6}$$

The overall reaction to synthesize the syngas to DME route is represented by the combination of reactions (4), (5) and (6) into reaction (7):

$$366 \qquad 3H_2 + 3CO \to CH_3OCH_3 + CO_2 \quad -\Delta H_{298 \ K; 0:1 \ MPa} = 246.0 \ kJ \ / \ mol \tag{7}$$

The overall reaction is exothermic and generates two molecules of products from six molecules of reactants. Hence, according to the Le Châtelier principle [76], conversion is favored working at high pressure and low temperature.

- 370 *3.2 Reaction kinetics*
- 371

The DME reactor was simulated in Aspen Plus with an RPLUG reactor combined with a Langmuir-Hinshelwood Hougen-Watson (LHHW) kinetic model based on three simultaneous reactions (Eqs.4,5,6). Bi-functional catalyst Cu/ZnO/Al₂O₃: γ -Al₂O₃ with a loading ratio of 1:2 has been selected from the literature, with the Cu/ZnO/Al₂O₃ active for the methanol synthesis, while

³⁴⁹

the γ -Al₂O₃ component catalyzes the methanol dehydration [72] among the other presented in the literature the selected is most investigated [77]. The details of the catalyst properties are reported in Table 2. The kinetic model adopted in the present simulations has been extensive used in the literature [14,71,78–80]. The rate expression for CO₂ hydrogenation, RWGS, methanol dehydration is given by equations (8-10) [72,81,82].

381
$$r_{CO_{2} hydrogenation} = \frac{k_{1} \left(p_{H_{2}} \cdot p_{CO_{2}} \right) \left[1 - \left(\frac{1}{K_{eq,1}} \right) \frac{p_{CH_{3}OH} \cdot p_{H_{2}O}}{p_{CO_{2}} p_{H_{2}}^{3}} \right]}{\left(1 + k_{2} \frac{p_{H_{2}O}}{p_{H_{2}}} + \sqrt{k_{3} \cdot p_{H_{2}}} + k_{4} \cdot p_{H_{2}O} \right)^{3}}$$
(8)

382
$$r_{RWGS} = \frac{k_5 \cdot p_{CO_2} \left[1 - \frac{1}{K_{eq,2}} \frac{p_{CO} \cdot p_{H_2O}}{p_{CO_2} \cdot p_{H_2}} \right]}{1 + k_2 \frac{p_{H_2O}}{p_{H_2}} + \sqrt{k_3 \cdot p_{H_2}} + k_4 \cdot p_{H_2O}}$$
(9)

383
$$r_{MeOH \ dehydration} = \frac{k_6 \cdot K_{CH_3OH}^2 \left[[\Pi]_{CH_3OH}^2 - \left(\Pi_{H_2O} \cdot \frac{\Pi_{DME}}{K_{eq,3}} \right) \right]}{\left(1 + 2\sqrt{K_{CH_3OH} \cdot \Pi_{CH_3OH}} + K_{H_2O} \cdot \Pi_{H_2O} \right)^4}$$
(10)

Reaction rates of equation (8-10) are expressed in kmol/kg_{cat} s, p is the partial pressure of the gases in Pa and Π the concentration expressed in kmol/m³. The equilibrium constant (K_i) and constant rate (k_i) values used to determine the reaction rates are shown in Table 4.

387

Table 4. Kinetic parameters used in DME synthesis.

	Pre	unit	B	unit
		unit	Ъ	unit
\mathbf{k}_1	1.07×10^{-13}	(kmol/(kg-sPa ²))	36,696	(J/mol)
\mathbf{k}_2	3450	-	0	(J/mol)
$k_{3}^{0.5}$	1.578×10 ⁻³	Pa ^{-0.5}	17,197	(J/mol)
k_4	6.62×10 ⁻¹⁶	Pa ⁻¹	124,119	(J/mol)
k_5	122	(kmol/(kg s Pa))	-94,765	(J/mol)
k ₆	1.486×10^{11}	(kmol/(kg s))	-143,666	(J/mol)
K _{CH3OH}	5.39×10 ⁻⁴	m ³ /kmol	70,560.92	(J/mol)
K _{H2O}	8.47×10 ⁻²	m ³ /kmol	42,151.98	(J/mol)

388

389 These parameters refer to the Arrhenius equation shown by equation (11).

390
$$k_i = (\operatorname{Pre})_i \times \exp\left(\frac{B_i}{RT}\right)$$
 (11)

where B represents either the activation energy or the reaction enthalpy or a combination of both [81] as in LHHW kinetic mechanism the rate constants are represented as combination of rate constants and equilibrium constants. The following expressions were used to determine the equilibrium constants [55,81,83].

395
$$\log_{10} K_{eq,1} = \frac{3066}{T} - 10.592$$
 (12)

396
$$\log_{10}(1/K_{eq_2}) = -\frac{2073}{T} + 2.029$$
 (13)

397
$$\ln K_{eq,3} = \frac{3220}{T} - 1.7$$
 (14)

The model was used to perform a sensitivity analysis of methanol and DME yield using the equations (15) and (16) varying the composition of the inlet stream, H₂:CO ratio, and the amount of the diluent H₂O and CO₂.

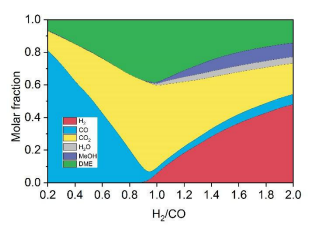
401
$$DME_{yield} = \frac{DME_{out}}{(CO + CO_2)_{in}}$$
 (15)

402
$$MeOH_{yield} = \frac{MeOH_{out}}{(CO + CO_2)_{in}}$$
 (16)

Where DME_{out} and $MeOH_{out}$ are the DME and methanol molar flow at the outlet of the reactor (stream 15) and CO and CO₂ the molar flow at the inlet (stream 14).

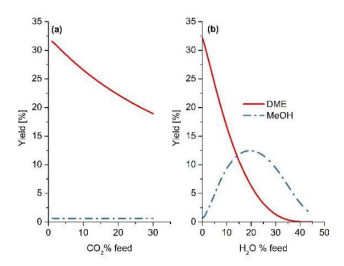
As shown in Figure 5 the highest DME yield is obtained by feeding a syngas with an $H_2/CO = 1$ 405 with a positive effect of having the main byproduct of the reaction as CO₂ which can be easily 406 separated from the DME and MeOH in the separation unit which is verified from the results 407 reported by Ogawa et al [41]. At H₂/CO=1, the DME yield was 38.6% and MeOH yield was 0.8% 408 which are similar to the results reported by Pozzo et al [71]. It is observed that with increasing the 409 CO₂ content at the inlet feed, the DME yield decreases. This is attributed mainly to two factors. 410 411 Firstly, the methanol synthesis is retarded with the increase of CO₂ content [84] as CO₂ molecules are absorbed by the methanol catalyst by occupying the active sites quicker than CO and H_2 . 412 affecting the MeOH production and consequently also the DME synthesis [85] as shown in Figure 413 6(a). Secondly, with a high CO₂ concentration in the feed of the DME reactor, the beneficial effect 414 of the water gas shift reaction would get decreased. The water formed is removed by WGS 415 producing hydrogen which kinetically advances the methanol production. Therefore, the higher CO₂ 416 favors the reverse-water gas shift that reduces the hydrogen content and produces more water. The 417 effect of higher water content at the inlet is even worse than CO_2 and it can be seen in Figure 6(b). 418 The high water percentage shifts the methanol dehydration towards the reactants, increasing the 419

420 MeOH yield while reducing the DME yield. With a water percentage higher than 20%, also 421 methanol production is penalized. In addition, the water tends to deposit near the catalyst 422 accelerating the catalyst degradation [41]. Therefore, to increase the DME production it is necessary 423 to have at the inlet of the DME reactor a syngas composed by an equimolar H₂-CO mixture, reduce 424 the CO₂ percentage (molar fraction) in the 0-5% range and remove as much as possible the water 425 content.



426

427 Figure 5: Influence of the H_2 /CO ratio on the equilibrium synthesis of DME at T=250°C and p=50 bar.



428

429 Figure 6: a) Influence of the CO_2 and b) H_2O on the equilibrium synthesis of DME at T=250°C and p=50 bar.

430 **4. Results**

431 **4.1 Effect of operating conditions**

432 A sensitivity analysis of the most influential parameters – namely: the operating pressure of the 433 chemical looping CO_2/H_2O splitting unit, outlet temperature of reduction reactor, CO_2/H_2O 434 composition in the oxidation reactor of the CL unit, and turbine inlet temperature – was performed 435 to maximize the global efficiency of the plant (Eq. 17) and the DME production.

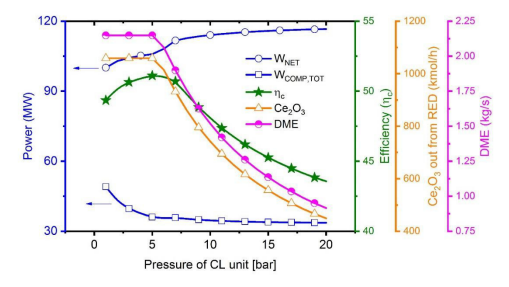
436
$$\eta_c = \frac{\dot{m}_{DME} \cdot LHV_{DME} + \dot{m}_{MEOH} \cdot LHV_{MEOH} + W_{NET}}{\dot{m}_{NG} \cdot LHV_{NG}}$$
(17)

15

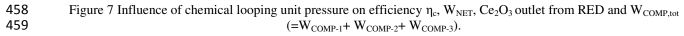
437 where: \dot{m}_{DME} , \dot{m}_{MeOH} represent the DME and MeOH streams produced (kg/s), while LHV_{DME}, 438 LHV_{MeOH}, and LHV_{NG} are the lower heating value (MJ/kg) of DME, MeOH and natural gas 439 respectively, and W_{NET} is the net power (MW) produced inside the plant with \dot{m}_{NG} being the natural 440 gas stream feed into the plant (kg/s).

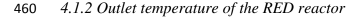
441 *4.1.1 Chemical looping (CO₂/H₂O) splitting (CL) unit pressure*

Figure 7 shows the effect of varying the pressure of the chemical looping unit, where both 442 oxidation and reduction reactors work at the same pressure. With the increase of pressure of the CL 443 unit, an efficiency gain is observed from 49.4% at 1 bar to 51.1% at 5 bar. This can be attributed to 444 the fact that a significant saving of the auxiliary power compression (W_{COMP,tot} in Figure 7) is 445 obtained by reducing the pressure ratio of syngas compression. However, with a further increment 446 of the pressure, the efficiency decreases, dropping down to 43.6% with 20 bar of pressure. Based on 447 the Le Châtelier principle, it can be understood that the reaction in the RED reactor is not 448 449 thermodynamically favored at high pressure since the reduction reaction has three moles of reactants and four moles of products. In fact, it can be seen that over 5 bar the amount of reduced 450 ceria (Ce_2O_3 line in Figure 7) at the outlet of the reactor decreases. This results in a lower syngas 451 452 production from the OXI reactor, as less reduced ceria is available, subsequently, the DME production drops effecting the overall plant efficiency. DME production drops after 5 bar pressure 453 and it does not vary between 1 to 5 bar while the W_{NET} increases very slowly from 100 to 105 MW. 454 The CL unit pressure can be fixed to 2 bar as the benefit of working at higher pressure is offset by 455 the power required to maintain pressure drop while working with solids. 456





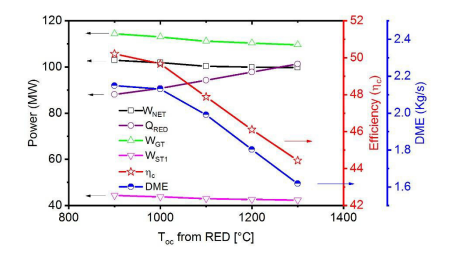




Another fundamental parameter is the outlet temperature of the OC from the RED. It is found that below 900°C there is no complete metal oxide conversion, similar to the results observed by Warren and Scheffe [31]. Therefore, all the analysis were performed considering reduction temperature above 900°C. A higher OC temperature at the outlet of the reactor, inherently demands

more heat supply. Since this heat is derived from the heat of combustion, to have higher RED 465 temperature, less CO₂ needs to be recirculated to the combustion chamber. This, even though results 466 in a corresponding drop in the power spent for recycling CO₂, also implies a lower mass flow 467 through the GT, producing less power, as shown in Figure 8. Such would then lower the power 468 produced by the ST1 as well, notwithstanding the higher temperature of the GT outlet, and hence 469 470 decreasing the net power output. In addition, a higher outlet temperature of RED also restricts the effective operation of the OXI. In fact, since both the CO₂ and H₂O splitting reactions are 471 exothermic, by principle, this requires the reactions to take place at a lower temperature. Moreover, 472 the water-splitting reaction has a higher exothermicity than CO₂ splitting with Ce₂O₃; thus, a higher 473 temperature would result in a slower reaction rate for H₂O splitting, resulting in a CO-rich syngas. 474 475 This effect is evident in Figure 8, in which a significant drop in the DME production can be seen beyond 1000°C (from 2.15 kg/s for 900°C to 2.13 kg/s at 1000°C and to 1.99 kg/s at 1100°C) due to 476 a deviation from the ideal H₂-CO ratio and higher concentration of CO₂ in the produced syngas 477 stream (Figure 9). 478

DME is one of the primary products of the proposed polygeneration system, a decrease in the DME production has a dramatic impact on the plant efficiency, as clearly observed in Figure 8 where the drop in the DME yield drives the trend of the decrease in the overall plant efficiency. To be more specific, a relative drop of 10.5% in efficiency is observed between 1000°C and 1300°C, corresponding to an in DME production of 24% and a relative net power output drop of 2%.



484

Figure 8 Influence of the metal oxide outlet temperature of RED on efficiency (η_c), W_{NET}, W_{GT} Q_{RED}, and DME production

487 Figure 9 shows the effect of the metal oxide inlet temperature to the OXI on the H_2/CO ratio in the syngas and on the CO₂ content in the syngas after water removal. As mentioned before, water 488 489 splitting is favored at a lower temperature compared to CO₂ splitting due to the higher exothermicity of the reaction. Hence, in order to have the ideal $H_2/CO = 1$ for DME production, it is 490 necessary to send an increasingly H₂O rich mixture with the higher temperature to the OXI. In fact, 491 with the increase of the OC temperature, the water splitting is further penalized and consequently, a 492 higher H₂O content ranging from 60% to 74% for OC temperature from 900-1300°C. In addition, as 493 494 already explained in section 2.2.5, the dilution of syngas with CO_2 has to be avoided in order to enhance DME production. As shown in Figure 9(b), even if it might be possible to produce the ideal 495 composition of syngas (i.e., 1:1 H₂/CO ratio) for any metal oxide temperature inlet, the CO₂ content 496

increases at higher temperatures. For this reason, it is suitable to work with lower ceria inlet temperature (900-1000°C) to avoid CO_2 dilution.

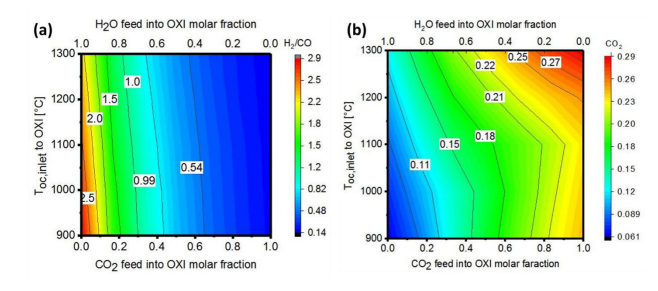


Figure 9 Effect of the gas mixture composition fed into OXI on a) final syngas H₂/CO ratio, b) CO₂ content (molar fraction) in the syngas after water removal.

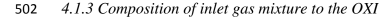
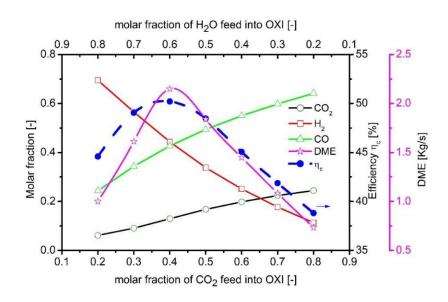
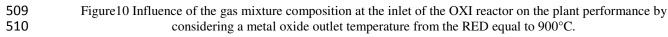


Figure 10 describes the effect of the variation of the gas mixture composition at the inlet of the OXI on plant performance. The maximum efficiency of 50% is achieved with an OXI inlet mixture of 60% of H₂O and 40% CO₂. In such a condition, the OXI outlet gas has the equimolar H₂:CO ratio (i.e., H₂ and CO curves intersect) which reflects the maximum DME production. Similar claims has been presented by Ohno et al [86].



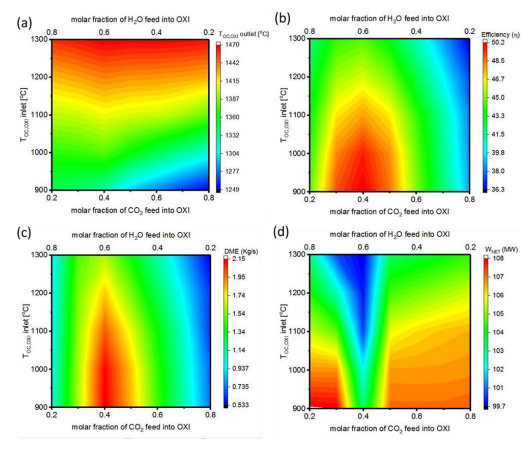
508

499



511 Increasing the water content in the feed to OXI reactor increases outlet metal oxide temperature 512 (Figure 11(a)) as water splitting is more exothermic than CO_2 splitting reaction. In the proposed

OXYF-CL-PFG plant layout, the oxidized ceria is recirculated back to the reduction reactor without 513 intermediate heat recuperation. Hence, a higher temperature of oxidized ceria at the outlet of OXI 514 results in a higher inlet temperature of OC to the RED which thereby reduces the heat requirement 515 for the reduction reaction. Due to inlet higher temperature of OC to the RED reactor, the heat 516 requirement from the combustion chamber reduces and therefore, the recirculation of CO₂ to the 517 518 combustion chamber to maintain the temperature of the outlet would be increased. With this, the power output from the gas turbine (GT) increases as higher flow expands which increases the net 519 power production, as seen in Figure 11(d). As stated earlier, this can be possible with the higher 520 H_2O concentration in the feed of OXI which increases the H_2/CO ratio more than unity leading to 521 the drop in DME production and overall efficiency (see Figure 11(b) and (c)). Therefore, an ideal 522 523 H₂/CO ratio feed to DME reactor, even though leads to lower overall net power, however, ensures the highest efficiency of the polygeneration unit, as can be understood from Figure 11(d). In the 524 case of a non-ideal H₂/CO ratio being fed to the DME reactor, it leads to a lower conversion with 525 unreacted syngas in the product stream. Even though after distillation, this is recycled to the 526 527 combustor increasing the power, but reduces the DME production and thus, the overall efficiency of 528 the plant.



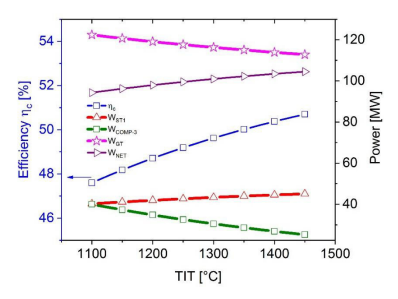
529 530

531

Figure 11 Effect of the gas mixture inlet composition and metal oxide inlet temperature (T_{OC,OXI} inlet) on (a) the temperature of the metal oxide outlet, (b) plant efficiency, (c) DME production, and (d) net power.

532 *4.1.4 Gas turbine inlet temperature*

533 Finally, the impact of the gas turbine inlet temperature (TIT) was analyzed. It can be said from 534 Figure 12 that with an increase of TIT the efficiency and net power produced are positively 535 influenced. Nevertheless, the output from the GT drops, which is the result of a lower recirculation of CO₂ to the combustor, needed to ensure a higher combustor exit temperature and consequently a higher TIT. This also causes a lower gas volume to be expanded within the GT, resulting in a lower power output, even though a partial compensation of the lost work is obtained by the lower compression work for the recirculated CO₂ in COMP-2. The power produced by the steam turbine ST1 increases slightly due to a higher exhaust temperature from the GT, overcoming the lower overall gas flow rate. For a TIT of 1100°C, the efficiency of 47.6% was obtained, which increases to 50.7% for a TIT of 1450°C.



543

544 Figure 12 Effect of the gas turbine inlet temperature TIT on the efficiency of the plant (η), power produced by the steam turbine 545 (W_{ST1}), by the gas turbine GT (W_{GT}) and the power absorbed by the COMP-3 (WCOMP-3)

546 **4.2 Plant performance**

- 547 Based on the sensitivity analysis the following operating parameters were chosen:
- 548 chemical looping operation pressure of 2 bar;
- 549 40% CO₂ and 60% H₂O in the gas mixture feed to the oxidation reactor with an excess of 550 60% with respect to the stoichiometry as per the Ce₂O₃ inlet to OXI;
- 551 reduction reactor outlet temperature 900°C;
- 552 TIT of 1377°C.

The plant performance based on the listed parameters is summarized in Table 5. Overall, the plant produces 102.90 MW_e, 185.6 ton/day (2.15 kg/s) of DME and 2.59 ton/day (0.03 kg/s) of methanol with a total efficiency of 50.21% and a DME yield of 24.9% (as per equation (8)). The highest power consumption is represented by the COMP-3 for the recycle of the CO₂ followed by the compression work in the ASU which accounts for 17% and 11.5% of the gross power generated respectively. Table 6 lists the composition and main thermodynamics parameters of major streams.

The inlet stream to the DME reactor (stream 14) has the ideal $H_2/CO = 1$ ratio, while the CO₂ content is 13%. However, it can be seen in Figure 11(b), that the minimum CO₂ percentage which can be achieved from the oxidation reactor is near 6%, even though not producing the equimolar mixture of H_2/CO . Therefore, the actual plant configuration allows producing a syngas with a composition which diverges from the ideal H_2/CO ratio of syngas. A solution might be to propose two distinct oxidation reactors, one for the CO_2 splitting and another for the water-splitting. However, this will lead to two different oxidized metal oxide temperatures, complicating the system design dynamics and operations.

567 An encouraging result is that the proposed oxyfuel-NGCC cycle with the chemical looping and 568 DME unit permits to cut the efficiency penalty of CCS. In particular, compared with results from 569 literature [87], it is possible to achieve a gain of 4 percentage points with respect to a stand-alone 570 Oxyfuel-NGCC process¹.

To evaluate the overall plant performance with regards to CO_2 savings, besides calculating the overall CO_2 avoided, the plant carbon efficiency, C_{eff} provides an alternative measure as per the following equation (18) [78,88,89].

574
$$C_{eff} = \frac{\dot{m}_{C,DME} + \dot{m}_{C,MeOH}}{\dot{m}_{C,NG}},$$
 (18)

where $\dot{m}_{C,DME}$ and $\dot{m}_{C,NG}$ are respectively the mass flow of carbon contained in the product (DME) and the primary reactant (inlet NG stream) respectively. $\dot{m}_{C,MeOH}$, denoting the mass flow of carbon in methanol contributes an minor fraction to the overall carbon efficiency of the polygeneration system as well.

579 The total CO₂ produced in polygeneration plant is 3.36 million tons per year out of which 3.4% is converted to the DME (CO_{2.DME}) with an overall plant carbon efficiency of 22.25% (Table7). The 580 recirculation streams of CO₂ (CO_{2,REC}) in the combustor accounts for the 85% produced CO₂ from 581 582 the exhaust (2.86 million tons per year), while the one sent into OXI for dissociation is 6.54%. In addition, a polygeneration scheme ensures the ability to produce DME within the same system, thus 583 cutting emissions from stand-alone DME production. Conventional DME production via a stand-584 alone steam methane reforming process results in an equivalent CO₂ emission of 51.1 kgCO₂/GJ of 585 DME [53]. Therefore, an equivalent of an additional 85.65 kilotonne of CO₂ was saved by the 586 polygeneration scheme accounting of total 589.15 kilotonne of CO₂ avoided annually. More 587 detailed share of carbon capture and utilization is listed in table S2 in supplementary file. As for the 588 carbon efficiency, the CL unit pressure and increase in CO₂ content to the OXI form the most 589 negative impacts, as can be observed from Table 7. This can be concluded from the decrease in 590 DME production, indicating the contribution of different operating conditions to the overall optimal 591 operation of the polygeneration plant. 592

593 Table 5 Plant results with selected parameters.

NG feed	25.2 ton/h
W _{GROSS}	167.61 MW
W _{NET}	102.90 MW
η_c	50.21%
W _{COMP-1}	3.76 MW
W _{COMP-2}	10.67 MW

¹ considering 0.09 kWh/Nm³ energy requirement for CO₂ compression [113].

W _{COMP-3}	28.29 MW
W _{ASU}	19.34 MW
W _{GT}	114.42 MW
W _{ST1}	44.30 MW
W _{ST2}	2.96 MW
W _{TURBEXP}	4.37 MW
\dot{m}_{DME}	185.6 ton/day (2.15 kg/s)
m _{MeOH}	2.59 ton/day (0.03 kg/s)
$CO_{2,REC}$	85%
$CO_{2,DME}$	3.4%

594

595 Table 6. Thermodynamics properties and composition of selected streams.

2	1 1		1										
Stream	28	10	14	15	17	20	31	37	38	43	47	7	9
T (°C)	900	1322	200	250	46	43	1377	80	40	40	-41	900	1322
P (bar)	2	2	5	50	10	9	26	26	1	1	10	2	2
Mole flow (kmol/s)	1	0.47	0.34	0.15	0.05	0.04	3.67	2.44	0.19	0.28	0.09	0.59	0.29
Molar fraction													
H ₂	0.57	0.32	0.44	0.04	0	0	0	0	0	0	0.01	0	0
H ₂ O	0	0.28	0	0.01	0.03	0	0.22	0	0	0.99	0	0	0
CO_2	0	0.09	0.13	0.6	0	0	0.77	0.99	0.99	0	0.96	0	0
CO	0.29	0.30	0.43	0.03	0	0	0	0.01	0	0	0.03	0	0
CH_4	0.12	0	0	0	0	0	0	0	0	0	0	0	0
other gases*	0.02	0.01	0	0	0	0	0	0	0.005	0.01	0	0	0
MeOH	0	0	0	0.01	0.02	0.01	0	0	0	0	0	0	0
DME	0	0	0	0.31	0.95	0.99	0	0	0	0	0	0	0
CeO ₂	0	0	0	0	0	0	0	0	0	0	0	1	0
Ce ₂ O ₃	0	0	0	0	0	0	0	0	0	0	0	0	1

*other gases include N_2 and other trace gases of natural gas

⁵⁹⁸ Table 7. Overall plant carbon efficiency for specific plant operating parameters.

Plant Operation Conditions	CL Unit Pressure [bar]	Toc from RED [°C]	H ₂ O%:CO ₂ % feed to OXI	Gas Turbine TIT [°C]	Carbon Efficiency (<i>Ceff</i>) [%]
Ideal	2	900	40:60	1377	22.25%
High CL Unit Pressure	20	900	40:60	1377	9.48%
High Toc from RED	2	1300	40:60	1377	16.70%
High CO ₂ % in OXI gas Feed	2	900	80:20	1377	7.60%
High H ₂ O % in OXI gas Feed	2	900	20:80	1377	10.40%
High Gas Turbine TIT	2	900	40:60	1450	22.25%

599

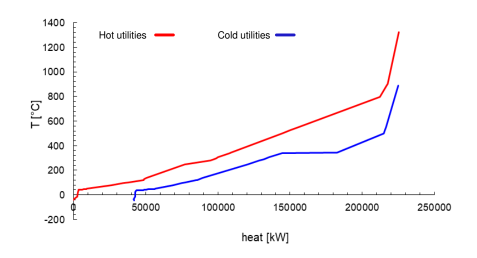
600 5. Pinch analysis

The thermal integration of the proposed polygeneration plant was performed using the pinch point analysis [90]. The highest temperature of 1322°C corresponds to the oxidation reactor outlet,

⁵⁹⁷

603 while the lowest of -40°C is the DME condensation temperature. Figure 13 shows the hot and cold 604 composite curve indicating a good thermal integration between cold and hot utilities, without the 605 use of an external heat source. Therefore, the scope for a further increment in the efficiency of the 606 power plant through optimized heat integration is limited. Starting from the hotter utilities, the 607 profile can be interpreted as follow:

- The cold utility curve from 50°C to 550°C represents the steam generation (stream 2A-5A and 2B-5B) driven by the exhaust gas from GT (stream 32) and the hot syngas from the oxidation (stream 10). It also represents the CO₂-H₂O preheating before the dissociation (stream 39 and 45) driven by stream 10a and 10b (hot syngas) in the HX-5 and HX-10 (for CO₂ preheating) and HX-4 and HX-12 (for H₂O preheating);
- 613 The steep part of the curves, from 550° to 900°C for the cold utilities, represents the of
 614 natural gas preheating before the reduction in RED (stream 5) taking place in the heat
 615 exchangers (HX-1 and HX-8B) and (HX-2 and HX-8);
- The part of the curves near and below zero is mostly related to the distillation unit and the
 condensation up to a temperature of -40°C of the DME;



618

619

Figure 13 Pinch point analysis with hot and cold composite curves

620

621 **6.** Exergy analysis

Exergy of a steady stream of matter is defined as the maximum amount of work obtainable when the stream is brought from its initial state to the dead state by processing during which the stream may interact only with the environment [91]. Exergy (E) can be divided into different components: kinetic exergy E_k , potential exergy E_{pot} , physical exergy E_{ph} and chemical exergy E_{ch} .

626
$$E = E_k + E_{pot} + E_{ph} + E_{ch}$$
 (19)

In the presented exergy analysis, the physical and chemical exergy are considered. The physical exergy is defined as the maximum work achievable from a system that from its initial state is

brought to the environmental state with only thermal and mechanical interaction with the 629 environment. While the chemical exergy is the maximum work obtainable from a system that is 630 brought from the environmental state to the dead state involving heat transfer and exchange of 631 substances only with the environment. The two types of exergy are given by equation (20) and 632 (21)[92]. In particular, for a mixture, the total chemical exergy $E_{ch.tot}$ is made by two contributes: 633 634 the chemical exergy of the single i-th component $E_{0,i}$ and the work obtainable from a reversible 635 isothermal expansion at T₀ from the partial pressure p₀₀ of the i-th component and environment pressure p_0 . 636

637
$$E_{ph} = \sum_{i} x_{i} \cdot \left[\left(h_{i} - h_{0,i} \right) - T_{0} \cdot \left(s_{i} - s_{0,i} \right) \right]$$
(20)

$$638 \qquad E_{ch,tot} = \sum_{i} x_i \cdot \left[E_{0,i} + \varphi \cdot R \cdot T_0 \ln \frac{p_0}{p_{00}} \right]$$
(21)

639 The exergy analysis is based on the second principle of thermodynamics, thus permits to 640 evaluate the so-called "destroyed" exergy ($I_{destroyed}$). Destroyed exergy represents the real loss in 641 the quality of energy that cannot be identified by means of a simple energy balance because the 642 conservation of energy will always apply. The following equation represents an exergy balance of a 643 general device in steady state condition:

644
$$\sum_{i} (E_{out,i} - E_{in,i}) = \sum_{i} W_i + \sum_{i} Q_i \left(1 - \frac{T_0}{T_i} \right) + I_{destroyed}$$
(21)

In equation (21): i) the members at the left side represent the exergy contribution of the $E_{in,i}$ inlet and $E_{out,i}$ outlet mass flows, ii) W_i represents the absorbed/produced work by the device, iii) the second member at the right side is the contribution from the heat exchanged, which represents the work obtainable from the heat flux Q_i operating with a Carnot machine, and iv) $I_{destroyed}$ is the irreversibility generated.

In order to estimate the exergy efficiency (or efficiency of the second principle) of a system is necessary to define the resource exergy flow of the process (Fuel) and product of the process (Product). The exergy efficiency is shown by the following equation:

$$\theta 53 \qquad \eta_{ex} = \frac{E_P}{E_F} \tag{23}$$

654 Where E_P represents the exergy of the product streams and E_F the exergy of the resource streams. 655 However, the only exergy efficiency does not give a complete framework of the plant or subsystem. 656 For this reason, an additional exergetic factor and other parameters were adopted [93]:

657 – Relative irreversibilities:
$$\chi_i = \frac{I_{i,destroyed}}{I_{tot,destroyed}}$$
 (24)

24

658 - Fuel depletion rate:
$$\theta_i = \frac{I_{i,destroyed}}{E_{F,plant}}$$
 (25)

659 – Productivity lack:
$$\xi_i = \frac{I_{i,destroyed}}{E_{P,plant}}$$
 (26)

660 - Exergetic factor:
$$\psi_i = \frac{E_{F,i}}{E_{F,plant}}$$
 (27)

661 A reference state was selected for the analysis (Table 8). For the environmental state, a pressure 662 (P_0) of 1 atm and a temperature (T_0) of 20°C were selected, while for the dead state the reference 663 environment of Szargut [94] was chosen.

Table 8 Environment state and dead state data of chemical exergy.

Dead State											
Chemical exergy E _{ch} (kJ/mol)											
H_2	СО	CO_2	H_2O_{vap}	H_2O_{liq}	N_2	CH_4	O_2	CeO ₂	Ce ₂ O ₃ [12]	DME[42]	MeOH[42
236.09	275.1	19.20	9.181	0.87	0.696	853.36	3.837	33.8	384.7	1414.5	715.52

665

666 Since in the proposed layout there are several chemical reactions, which change the 667 composition of the gaseous streams, the first step was to evaluate the reference chemical exergy of 668 the multiple mixture streams using the dead state of the reference elements. The results are shown 669 in Table 9.

670	Table 9. Specif	fic chemic	al exergy of th	e gas mixture stre	eams.		
	Stream	31	28	10	15	13	47
	e _{ch,i} [kJ/kg]	389	27110	7391	11287	11919	6225

671

The exergetic performance of the overall plant has been assessed by evaluating its exergetic efficiency (Eq. 28) and the total irreversibility generated (Eq. 29). As can be clearly observed pinch analysis, due to both electricity and heat self-sufficiency of the system, the input fuel, namely natural gas, contributes entirely to the net exergy input to the system (i.e., it is 100% of the total exergy input). The products are the total DME, MeOH and the net power produced by the plant.

677
$$\eta_{ex} = \frac{\dot{m}_{DME} \cdot E_{DME} + \dot{m}_{MEOH} \cdot E_{MEOH} + W_{NET}}{\dot{m}_{CH_4} \cdot E_{NG}}$$
(28)

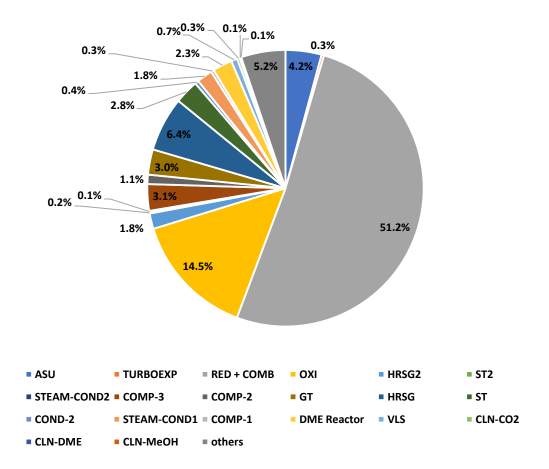
678
$$I_{tot, destroyed} = \dot{m}_{CH_4} \cdot E_{NG} - \dot{m}_{DME} \cdot E_{DME} - \dot{m}_{MEOH} \cdot E_{MEOH} - W_{NET}$$
(29)

As expected, since the exergy efficiency is primarily influenced by to the fuel value of natural gas,
DME, MeOH, and the net power produced, the exergy efficiency trend is specular to the

thermodynamic efficiency previously described. The energetic and exergetic efficiency with respect to the pressure of CL unit and metal oxide inlet temperature to reduction reactor and turbine inlet temperature is presented in Figure S1 and effect of the molar composition of CO_2 and H_2O in OXI is presented in Figure S2 in supplementary file. Finally, also a detailed exergy analysis of the components of the layout operating at the conditions described in section 4 was performed. Chemical, physical and total exergy values of all streams are reported in Table S3 in supplementary file.

The global exergy efficiency (η_{ex}) of the plant was obtained as 45.0%, five points lower than the calculated first-law efficiency. The total irreversibilities generated are 202.72 MW with an overall fuel depletion rate (θ) of 53.84%. All the components present an exergetic efficiency over the 80%, with the exception of the ASU (55.9%), the two condensers for the steam cycle (32%) and the CLN-MeOH (77%). However, the contribution of COND-A, COND-B, and ASU to the overall efficiency is marginal since their relative irreversibilities χ_i do not exceed the 3.9% (see Table 10).

The exergy efficiency of the RED+COMB results in 88.1%. Although this value is not 694 extremely low, more than half of the 202.72 MW total irreversibilities are in this component (Figure 695 14). As shown in Table 10, the RED+COMB exergetic factor ψ results in 231.3%, so the 696 irreversibilities are not due to the efficiency but are mainly proportionally correlated to the high 697 698 exergy of the inlet streams. In fact, the exergy inlet of the RED+COMB ranks first among the components (870 MW), the second being the turbo-expander inlet (378 MW). Moreover, it is worth 699 mentioning that, with the hypothesis of zero heat losses inside the combustor and reduction reactor, 700 the main contributor to the exergy losses are of the chemical form. In fact, the exergy efficiency of 701 the RED+COMB, considering only the chemical exergy of the inlet and outlet streams, results in 702 70%. The oxidation reactor is the second-ranked component for the relative irreversibilities 703 704 parameter (14.4%) even if the exergy efficiency (83.4%) results in to be lower than the one of the RED+COMB. This is due to the lower exergy factor (50.7%). 705



706

707

Figure 5 Total irreversibilities distribution.

The other irreversibilities are mostly from the HRSG1 of the steam ranking cycle (SRC1) (13.99 MW) and in the compression process (9.3 MW). The DME reactor jacketing for saturated steam production allows increasing the exergy efficiency of the component of 2.2%.

711 Table 10 Results from the exergetic analysis of the main components.

Componet	Exergy balance eq. ¹	I [MW]	η _{ex,i} [%]	$\begin{array}{c} \theta_{i} \\ [\%] \end{array}$	Ψi [%]	ξ _i [%]
ASU	$\mathbf{E}_0 + \mathbf{W}_{\mathrm{ASU}} = \mathbf{E}_1 + \mathbf{E}_{\mathrm{1b}} + \mathbf{I}_{\mathrm{ASU}}$	8.53	55.91	2.26	5.14	5.03
TURBOEXP	$E_3 = W_{TURBOEXP} + I_{TURBOEXP}$	0.59	99.84	0.16	100.41	0.35
RED+COMB		103.83	88.08	27.58	231.29	61.23
OXI	$E_7 + E_{46} + E_{40} = E_{10} + E_{9a} + I_{OXI}$	29.47	84.57	7.83	50.74	17.38
HRSG2	$E_{10} + E_{2-B} = E_{10a} + E_{5-B} + I_{HSRG-2}$	3.68	97.49	0.98	38.88	2.17
ST2	$E_{5-B} = E_{6-B} + W_{ST2} + I_{ST2}$	0.38	89.77	0.10	0.99	0.22
COND-B	$E_{6-B} = E_{7-B} + I_{COND-B}$	0.25	32.0	0.07	0.10	0.15
COMP-4	$E_{38} + W_{COMP-4} = E_{39} + I_{COMP-4}$	0.08	98.26	0.02	1.18	0.04
COMP-3	$E_{49} + W_{COMP-3} = E_{37} + I_{COMP-3}$	6.29	91.73	1.67	20.21	3.71
COMP-2	$E_{29} + W_{COMP-2} = E_{30} + I_{COMP-2}$	2.21	99.33	0.59	87.57	1.30
GT	$E_{31} = W_{GT} + E_{32} + I_{GT}$	6.00	97.57	1.59	65.57	3.54
HRSG1	$E_{31} + E_{2-A} = E_{33} + E_{5-1} + I_{HSRG-1}$	12.88	90.0	3.42	34.21	7.60
ST1	$E_{5-A} = E_{6-A} + W_{ST1} + I_{ST1}$	5.68	89.75	1.51	14.72	3.35
COND-2	$E_{33} = E_{34} + E_{41} + I_{COND-2}$	0.88	98.51	0.23	15.80	0.52

COND-A	$E_{6-1} = E_{7-A} + I_{COND-A}$	3.71	32.00	0.99	1.45	2.19
COMP-1	$E_{13} + W_{COMP-1} = E_{14} + I_{COMP-1}$	0.71	99.10	0.19	20.88	0.42
DME Reactor	$E_{14} + E_{44} = E_{15} + E_{45} + I_{DME \ reactor}$	4.64	94.05	1.23	20.73	2.74
VLS	$E_{15} = E_{47} + E_{17} + I_{VLS}$	1.45	97.99	0.39	19.16	0.86
CLN-CO ₂	$E_{19} + Q^*_{\text{COND, CLN-CO2}} + Q^*_{\text{RED, CLN-CO2}} = E_{20} + E_{21} + I_{\text{CLN-CO2}}$	0.60	99.15	0.16	18.56	0.35
CLN-DME	$\begin{split} E_{22} + Q^*_{\text{COND,CLN-DME}} + Q^*_{\text{REB,CLN-DME}} \\ = E_{22} + E_{23} + I_{\text{REB-DME}} \end{split}$	0.30	99.56	0.08	17.90	0.18
CLN-MeOH	$E_{25} + Q^*_{\text{COND,CLN-MeOH}} + Q^*_{\text{REB,CLN-MeOH}}$ $= E_{22} + E_{23} + I_{\text{REB-MeOH}}$	0.15	77.39	0.04	0.18	0.09
NG _{PHX1}	$E_2 + E_{28} = E_3 + E_{28b} + I_{NG-PHX1}$	0.41	99.94	0.11	185.39	0.24
NG _{PHX2}	$E_4 + E_{28b} = E_5 + E_{29} + I_{NG-PHX2}$	1.59	99.78	0.42	188.17	0.94
CO _{2PHX}	$E_{10a} + E_{39} = E_{10b} + E_{40} + I_{CO2-PHX}$	0.38	99.48	0.10	19.34	0.23
H_2O_{PHX}	$E_{10b} + E_{45} = E_{11} + E_{46} + I_{H2O\text{-}PHX}$	3.26	95.63	0.87	19.80	1.92

712

¹The left-side of the equation in the table represents the fuel of the component, while the right side

of the equation represents the product and the irreversibility of the component.

715 Q^* represents the exergy obtainable using the heat of the selected stream

717
$$Q_{\text{COND,CLN-CO2}}^* Q_{\text{COND,CLN-CO2}} \times \left(1 - \frac{T_{\text{COND,CLN-CO2}}}{T_0}\right);$$
718
$$Q_{\text{REB,CLN-CO2}}^* = Q_{\text{REB,CLN-CO2}} \times \left(1 - \frac{T_0}{T_{\text{REB,CLN-CO2}}}\right);$$
719
$$Q_{\text{COND,CLN-DME}}^* Q_{\text{COND,CLN-DME}} \times \left(1 - \frac{T_0}{T_{\text{COND,CLN-DME}}}\right);$$
720
$$Q_{\text{REB,CLN-DME}}^* = Q_{\text{REB,CLN-DME}} \times \left(1 - \frac{T_0}{T_{\text{REB,CLN-DME}}}\right);$$
721
$$Q_{\text{COND,CLN-MeOH}}^* = Q_{\text{COND,CLN-MeOH}} \times \left(1 - \frac{T_0}{T_{\text{COND,CLN-MeOH}}}\right);$$
722
$$Q_{\text{REB,CLN-MeOH}}^* = Q_{\text{REB,CLN-MeOH}} \times \left(1 - \frac{T_0}{T_{\text{REB,CLN-MeOH}}}\right);$$
723

723

724 **7. Economic analysis**

An economic assessment was performed to calculate the capital cost of investment for the 725 726 construction of the proposed plant. The National Energy Technology Laboratory (NETL) guidelines 727 for techno-economic analysis for power plants was adopted [95,96]. This methodology defines capital cost at five levels: bare erected cost (BEC), engineering, procurement, and construction cost 728 (EPCC), total project cost (TPC), total overnight cost (TOC) and total as-spent cost (TASC). In the 729 current study, the TOC was considered for the capital investment expenditure. The first four items 730 are "overnight cost" and are expressed in base-year US dollar that is the first year of capital 731 expenditure. The Bare Erected Cost (BEC) comprises the cost of the equipment, facilities and 732 infrastructure, and labor required for its installation. The equipment cost was estimated using the 733 scaling factor exponent M, as given by equation (30) [97] and details of which can be found in [12]. 734

735
$$C_{equ} = C_{equ,ref} \left(J/J_{ref} \right)^M$$
(30)

where C_{equ} and $C_{equ,ref}$ represent the equipment cost with a capacity of J and J_{ref} , respectively.

To assess further costs related to setting up of the polygeneration plant including installation and other direct and indirect costs related to the project development, a bottom-up approach following the methodology adopted in the CAESER project [98] was selected. All the estimated equipment costs (Table S5 in supplementary file) were converted to the year 2017 US dollar using the chemical engineering plant cost index (CEPCI, Table S6 in supplementary file).

742
$$C_{equ,actual} = C_{equ,ref} \frac{CEPCI 2017}{CEPCI at the time of original cost}$$
 (31)

The cost of the cooling tower system was included in the cost of the four condensers (COND-1, COND-2, SRC1, and SRC2 condenser). The overall cost was subdivided between the four components proportionally to the calculated rejected heat. The cost of the two condensers (COND-A and COND-B) of the two HRSG were included in the HRSG investment cost. The most expensive equipment is the ASU, followed by the GT. The RED+COMB unit accounts for 5.2% of the total expenditure. The individual contribution of the respective equipment to the total overnight cost is shown in Figure 15.

The bare erected cost (BEC) of each equipment was given summing all the installation costs (see Table S6 in the supplementary file for assumptions of CAPEX estimation) to the equipment cost is given by equation (32).

$$BEC = C_{equ,actual} + Installation Cost$$
(32)

The engineering, procurement and construction cost (EPCC) comprises the BEC plus the costs of all services provided by the engineering, procurement and construction contractor (equation (33)). These items include detailed design, contractor permitting and project management costs.

$$F757 EPCC = BEC + INDIRECT COST$$
(33)

The total project cost (TPC) also includes the contingencies cost (equation (34))to account for unknown costs that are omitted or unheralded due to lack of complete project definition or uncertainties with the development status of a technology. In the present case, since the proposed plant is based on a novel technology arrangement, a high contingency cost of 30% was selected (Table S6 in supplementary file).

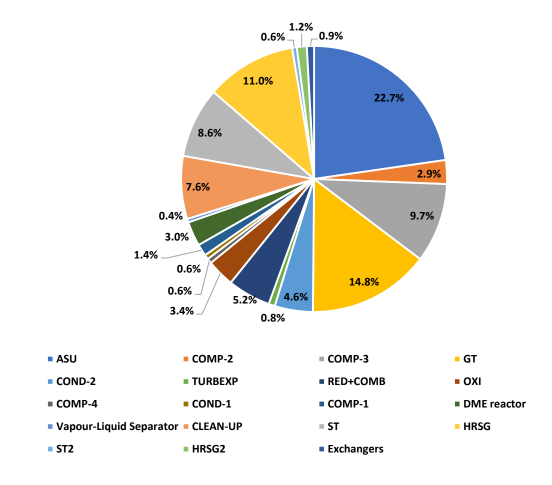
$$TPC = EPCC + CONTINGENCIES$$
(34)

The total overnight cost (TOC) comprises the TPC plus other overnight costs (Table S4 in supplementary file), owner's cost included (i.e pre-production, inventory capital, land, financing), it was calculated as:

$$TOC = TPC + OWNER'S COST$$
(35)

From the assumptions listed in Table S6 (in supplementary file), the total overnight cost (TOC) of the plant resulted in 537.45 \$million. Figure 15 represents the contribution to the total overnight

cost of the different equipment. The most expensive equipment resulted in the ASU, followed by
the GT. The RED+COMB unit accounted for 5.2% of the total expenditure. For the economic
analysis, all assumptions are listed in Table S7 and S8 in the supplementary file for OPEX
estimations.



774

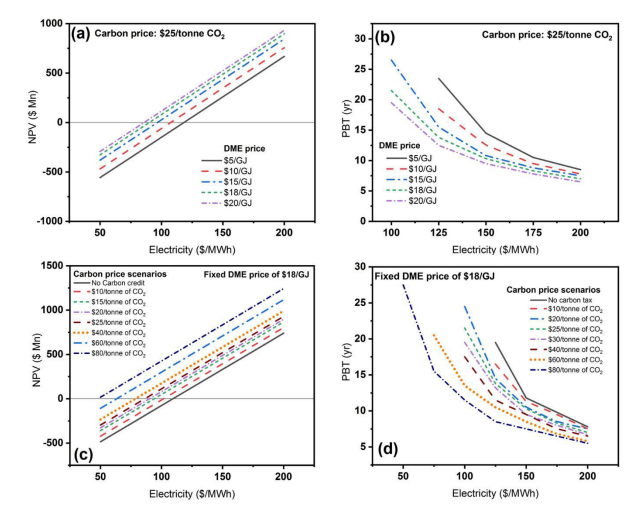
775

Figure 15 Contribution of the components to the TOC.

Finally, to evaluate the profitability of the plant during its lifetime a discounted cash flow analysis (DCF) was adopted. DCF is based on the concept of the time value of money, all the future cash flows are estimated and discounted by a discounted factor (i) (Table S8 in supplementary file), obtaining their present value [99]. The sum of the all discounted cash flows, both positive (revenues) and negative (operation cost, Table S7 in supplementary file), gives the net present values (NPV) as shown by equation (36).

782 NPV =
$$-\text{TASC} + \sum_{l=1}^{t} \frac{(\text{net cash flows})_l}{(1+i)^l}$$
 (36)

A project is acceptable only if the NPV is positive. TASC is used to evaluate the total project cost instead of TOC, in order to asses both escalation and interest during construction (Table S7 in supplementary file) [95,96]. A sensitivity analysis was performed to evaluate the effect of the selling price of power and DME on the economic performance of the plant.



787

Figure 16 (a) and (c) Economic performance varying carbon credits and DME prices for different electricity prices (b) and (d) Payback period (PBT) varying DME and electricity price for different levels of carbon credit

790	It is observed that a payback period (PBT) of 20 years was obtained with the electricity and the
791	selling price of DME of \$20/GJ and \$220/MWh respectively which is around 2.2 times the current
792	whole sale electricity price without carbon credits considered

793 Figure 16 (a) and (b) represents the variation of electricity prices for different DME prices from \$5-20/GJ. It is observed that for the current reported carbon credit price of \$25/tonne of CO₂ [100] 794 the NPV is positive if the electricity price of \$90/MWh (which is below current electricity price 795 with CCS) with a payback period of 21.5 years. Nonetheless, a rapid increase in the CO₂ price 796 between \$40 and \$80 per tonne is required to meet the agreements of Paris 2015 [101]. This trend 797 798 can be observed in many countries where high carbon tax of over 55\$/tonne prevails including Switzerland, Finland, Sweden, Liechtenstein, Norway, Mexico, Chile and many others [101]. 799 Indeed, such rapid rise in the carbon price have already started to materialize, with further increase 800 to be seen developed economies by 2030 [102–105]. Still many developing countries are struggling 801 to adopt the carbon price and emission trading system making it difficult to match the carbon price 802 difference. Therefore, the NPV values was varied for different scenarios of electricity and DME 803 price based on the carbon credit variation from \$10-80/tonne of CO₂ [106]. Figure 16 (c) and (d) 804 805 corresponds to the fixed current market DME price of \$18/GJ [53] and varying the carbon credits for electricity pricing. From Figure 16 (c) it can be said with the carbon price above \$40/tonne CO2 806 would have positive NPV that can be able to match with the current selling price of electricity is 807

above \$100/MWh with carbon capture [106–111] for the proposed polygeneration with a PBT of 17.5 years, with potential to drop further for higher carbon credit scenarios of \$60 and \$80/tonne of CO₂ the payback period reduces by nearly 2/3 (11.5 years).

However, more strong carbon credit policies and a further development of technologies, such as 811 Oxyfuel combustion, air separation, and chemical looping, will make the proposed polygeneration 812 813 plant more competitive. By considering oxygen transport reactors that use ion transport membranes, such as perovskites, for oxygen separation at high temperatures (i.e., above 700°C), high-purity 814 oxygen could be produced at a relatively lower price compared to ASU, thereby increasing the 815 efficiency and decreasing the equipment cost. At present, with this technology, it is possible to 816 produce 2000 tonne per day, which is sufficient for an oxyfuel plant of 110 MW capacity. 817 Therefore, with the adoption of the ion transport membrane technology that costs 31% less 818 compared to the ASU, consequently, the cost of DME and power production would decrease 819 tremendously and the overall efficiency of the plant would improve by 2-4% [112]. 820

821 8. Conclusion

822

A novel natural gas feed polygeneration plant was proposed that integrates a chemical looping 823 CO₂/H₂O splitting unit with an oxyfuel combustion unit for the production of power, DME and 824 methanol. The results demonstrated the advantages of using a chemical looping CO₂/H₂O process in 825 a polygeneration plant to reduce the efficiency penalty due to the carbon capture. The analysis 826 827 revealed that the ideal H₂/CO ratio for a single step DME synthesis to be which can be obtained by feeding H₂O/CO₂ ratio of 60/40%. The plant was able to produce 103 MWe and 185.6 ton/day (2.15 828 kg/s) of DME with an energetic and exergetic efficiency of 50.2% and 45.0%, respectively. 829 Compared to only power plant with carbon capture the present polygeneration revealed an 830 efficiency gain of 4%. Through an exergy analysis, the main contributors of exergy destruction 831 were identified: the combustor and reduction system resulted to contribute for 51.2% of the total 832 generated irreversibilities (221 MW). The capital investment was estimated to be \$534 million. The 833 overall CO₂ produced was 3.36 million tonne for 7446 hours (with a capacity factor of 0.85) of 834 835 annual operation of which approximately 3.4% is contributed by the DME production in a polygeneration scheme accounting for 589.15 kilotonne of total CO₂ avoided annually. Economic 836 analysis revealed that around 23% of the total equipment costs is attributed by ASU and with the 837 838 use of more sophisticated technology for producing oxygen at less price would decrease the capital 839 investment. A discounted cash flow analysis revealed that the proposed plant would able to meet the electricity and DME price of \$100/MWh an \$18/GJ with the carbon credit of \$40/tonne of CO₂, 840 which is projected to be the carbon credit by 2020. With stringent carbon pricing of \$80/tonne of 841 CO₂ the electricity price would drop below \$50/MWh. 842

843 Author Contributions:

- A Farooqui and F Di Tomaso developed the model and performed simulations. A Farooqui, F Di
- Tomaso, A Bose and D Ferrero written the manuscript in multiple iterations. J Llorca and M
- 846 Santarelli supervised the work and made the necessary modifications required in the manuscript.
- 847 The figures were made in multiple iterations by A Farooqui and Di Tomaso.

848	Glossary:	
849	ṁ	Mass flow rate (kg/s)
850	LHV	Lower heating value (MJ/kg)
851	W _{NET}	Net power produced inside the plant (MW)
852	W _{ST1}	Power produced in steam turbine of SRC1 (MW)
853	W _{ST2}	Power produced in steam turbine of SRC2 (MW)
854	W _{GT}	Power produced in gas turbine (MW)
855	W _{COMP,tot}	Auxiliary compression power in compressors (MW)
856	W _{COMP-3}	Auxiliary compression power for recycling CO_2 (MW)
857	CO _{2,DME}	CO ₂ embedded in DME (%)
858	$CO_{2,REC}$	CO_2 recycled within the plant (%)
859	E	Total exergy (MW)
860	E _k	Kinetic exergy
861	$E_{\rm ph}$	Physical exergy
862	E _{pot}	Potential exergy
863	E _{pot}	Chemical exergy
864	L _{ch}	Mass fraction (-)
865	h	Mass enthalpy (kJ/kg)
866	S	Mass entropy (kJ/kg-K)
867	φ	Activity coefficient for the i-th component
868	R	Ideal gas constant (8.314 J/mol-K)
869	ρ	Density (kg/m^3)
870	p ₀₀	Partial pressure of the i-th component (Pa)
871	W	Absorbed/produced work by the device (MW)
872	Q	Work obtained from heat flux (MW)
873 874	I _{destroyed} T	Irreversibility generated (MW) Temperature (°C)
874 875		Exergy efficiency (-)
876	η _{ex} n	Energy efficiency (-)
877	$\eta_c Q^*$	Exergy obtainable using the heat of the selected stream
878	Q C _{equ}	Equipment cost
879	C _{equ} ,actual	Actual component cost
880	C _{equ,ref}	Reference component cost
881	M	Scaling factor (-)
882	l	Number of years (-)
883	i	Discounted factor (-)
884	Р	Pressure (bar)
885	χ	Relative irreversibilities
886	θ	Fuel depletion rate (%)
887	بح	Productivity lack (%)
888	Ψ	Exergetic factor (%)
889	P ₀	Pressure at environment state of 1 atm
890	T_0 r^0	Temperature at environment state of 20°C
891	E^0	Standard chemical exergy
892 802	ΔG^0	Gibbs free energy
893 804	e _{ch}	Specific chemical exergy (kJ/kg)
894 805	K _i	equilibrium constant (m ³ /kmol) Arrhenius rate constant
895	k _i	Armenius rate constant

896	r	reaction rate (kmol/kg _{cat} s)
897	Π	concentration expressed (kmol/m ³)
898	В	activation energy (J/mol)
899	Pre	pre-exponential factor
900	RGIBBS	GIBBS Reactor where the calculations are based on minimizing the Gibbs
901	energy	
902	OC	Oxygen Carrier
903	C_{eff}	Plant carbon efficiency
904	-55	
905	Subscripts	
906	ph	physical
907	ch	chemical
908	vap	vapor state
909	liq	liquid state
910	COMP	compression work
911	TURBEXP	work by turbo expansion
912	0	environment state of 1 atm and 20°C
913	is	isentropic compressor
914	mech	mechanical
915	comp	compressor
916	turb	turbine
917	ex	exergy
918	in,i	contribution to inlet mass flows
919	out,i	contribution to outlet mass flows
920	tot	total or cumulative
921	i	i-th component in the mixture
922	COND	condenser
923	REB	reboiler
924	k	k-th component
925	Р	product stream
926	F	resource stream
927	Acronyms	
928		
929	NG	Natural Gas
930	VLS	Vapor Liquid Seperator
931	CLN	Column
932	SRC	Steam Rankine Cycles
933	MR	Methane Reduction
934	WS	Water Splitting
935	CDS	Carbon dioxide Splitting
936	DME	Dimethyl Ether
		•
937	ASU	Air Separation Unit
938	TURBOEXP	Turbo Expander
939	RED	Reduction Reactor
940	OXI	Oxidation Reactor
941	LHHW	Langmuir-Hinshelwood Hougen-Watson
942	ST	Steam Turbine
943	GT	Gas Turbine

944	PHX	Y Preheater
945	TPC	Total Project Cost
946	BEC	
947	EPC	
948	TOC	e
949	TAS	1
950	TPC	5
951	NET	L National Energy Technology Laboratories
952	CEP	CI Chemical Engineering Plant Cost Index
953	OPE	X Operational Expenditure
954	CAP	1 1
955	DCF	Discounted Cash Flow
956	NPV	Net Present Value
957	HR	Heat Rate
958	CSP	Concentrating Solar Power
959	CCS	Carbon Capture and Sequestration
960	PBT	Payback period in years
961		
962	REF	TERENCES:
963 964	[1]	I.E.A. IEA, CO2 Emissions from Fuel Combustion 2017 - Highlights, Int. Energy Agency. 1 (2017) 1–162. doi:10.1787/co2_fuel-2017-en.
965	[2]	IPCC, IPCC special report on global warming of 1.5oC, 2019.
966	[4]	http://www.ipcc.ch/report/sr15/.
967	[3]	P. Viebahn, D. Vallentin, S. Höller, Prospects of carbon capture and storage (CCS) in
967 968	[3]	China's power sector - An integrated assessment, Appl. Energy. 157 (2015) 229–244.
969		doi:10.1016/j.apenergy.2015.07.023.
	E 4 3	
970 071	[4]	M.E. Boot-Handford, J.C. Abanades, E.J. Anthony, M.J. Blunt, S. Brandani, N. Mac Dowell,
971 972		J.R. Fernández, MC. Ferrari, R. Gross, J.P. Hallett, R.S. Haszeldine, P. Heptonstall, A. Lyngfelt, Z. Makuch, E. Mangano, R.T.J. Porter, M. Pourkashanian, G.T. Rochelle, N. Shah,
972 973		J.G. Yao, P.S. Fennell, Carbon capture and storage update, Energy Environ. Sci. 7 (2014)
974		130–189. doi:10.1039/C3EE42350F.
	[5]	
975 976	[5]	R.P. Cabral, N. Mac Dowell, A novel methodological approach for achieving £/MWh cost reduction of CO2 capture and storage (CCS) processes, Appl. Energy. 205 (2017) 529–539.
970 977		doi:10.1016/j.apenergy.2017.08.003.
978	[6]	Y. Zhao, B. Jin, Z. Deng, Y. Huang, X. Luo, Z. Liang, Thermodynamic analysis of a new
979		chemical looping process for syngas production with simultaneous CO2capture and
980 081		utilization, Energy Convers. Manag. 171 (2018) 1685–1696.
981		doi:10.1016/j.enconman.2018.06.101.
982	[7]	Q. Zhang, Nurhayati, C.L. Cheng, D. Nagarajan, J.S. Chang, J. Hu, D.J. Lee, Carbon capture
983		and utilization of fermentation CO2: Integrated ethanol fermentation and succinic acid
984		production as an efficient platform, Appl. Energy. 206 (2017) 364–371.
985		doi:10.1016/j.apenergy.2017.08.193.
986	[8]	C. Graves, S.D. Ebbesen, M. Mogensen, K.S. Lackner, Sustainable hydrocarbon fuels by
987		recycling CO2 and H2O with renewable or nuclear energy, Renew. Sustain. Energy Rev. 15
		35

- 988 (2011) 1–23. doi:10.1016/j.rser.2010.07.014.
- [9] S.M. Jarvis, S. Samsatli, Technologies and infrastructures underpinning future CO2value
 chains: A comprehensive review and comparative analysis, Renew. Sustain. Energy Rev. 85
 (2018) 46–68. doi:10.1016/j.rser.2018.01.007.
- [10] A.E. Farooqui, M.A. Habib, H.M. Badr, R. Ben-Mansour, Modeling of ion transport reactor
 for oxy-fuel combustion, Int. J. Energy Res. 37 (2013). doi:10.1002/er.2923.
- R. Ben-Mansour, M.A. Habib, H.M. Badr, F. Azharuddin, M. Nemitallah, Characteristics of
 oxy-fuel combustion in an oxygen transport reactor, in: Energy and Fuels, ACS Publications,
 2012: pp. 4599–4606. doi:10.1021/ef300539c.
- A. Farooqui, A. Bose, D. Ferrero, J. Llorca, M. Santarelli, Techno-economic and exergetic
 assessment of an oxy-fuel power plant fueled by syngas produced by chemical looping CO2
 and H2O dissociation, J. CO2 Util. 27 (2018) 500–517. doi:10.1016/j.jcou.2018.09.001.
- [13] K. Jana, A. Ray, M.M. Majoumerd, M. Assadi, S. De, Polygeneration as a future sustainable energy solution – A comprehensive review, Appl. Energy. 202 (2017) 88–111.
 doi:10.1016/j.apenergy.2017.05.129.
- [14] Y.K. Salkuyeh, T.A. Adams, A new power, methanol, and DME polygeneration process
 using integrated chemical looping systems, Energy Convers. Manag. 88 (2014) 411–425.
 doi:10.1016/j.enconman.2014.08.039.
- [15] S. Li, L. Gao, H. Jin, Realizing low life cycle energy use and GHG emissions in coal based polygeneration with CO2 capture, Appl. Energy. 194 (2017) 161–171.
 doi:10.1016/j.apenergy.2017.03.021.
- 1009 [16] A. Bose, K. Jana, D. Mitra, S. De, Co-production of power and urea from coal with CO2 capture: Performance assessment, Clean Technol. Environ. Policy. 17 (2015) 1271–1280.
 1011 doi:10.1007/s10098-015-0960-7.
- 1012 [17] K. Jana, S. De, Environmental impact of biomass based polygeneration A case study
 1013 through life cycle assessment, Bioresour. Technol. 227 (2017) 256–265.
 1014 doi:10.1016/j.biortech.2016.12.067.
- 1015 [18] H. Huang, S. Yang, P. Cui, Design concept for coal-based polygeneration processes of
 1016 chemicals and power with the lowest energy consumption for CO2capture, Energy Convers.
 1017 Manag. 157 (2018) 186–194. doi:doi.org/10.1016/j.enconman.2017.11.073.
- 1018 [19] Y.K. Salkuyeh, T.A. Adams, A new power, methanol, and DME polygeneration process
 1019 using integrated chemical looping systems, Energy Convers. Manag. 88 (2014) 411–425.
 1020 doi:10.1016/j.enconman.2014.08.039.
- [20] R. Stanger, T. Wall, R. Spörl, M. Paneru, S. Grathwohl, M. Weidmann, G. Scheffknecht, D.
 McDonald, K. Myöhänen, J. Ritvanen, S. Rahiala, T. Hyppänen, J. Mletzko, A. Kather, S.
 Santos, Oxyfuel combustion for CO2capture in power plants, Int. J. Greenh. Gas Control. 40 (2015) 55–125. doi:10.1016/j.ijggc.2015.06.010.
- [21] V.N. Nguyen, L. Blum, Syngas and synfuels from H2O and CO2: Current status, Chemie Ingenieur-Technik. 87 (2015) 354–375. doi:10.1002/cite.201400090.
- 1027 [22] G.P. Smestad, A. Steinfeld, Review: Photochemical and thermochemical production of solar
 1028 fuels from H 2O and CO 2 using metal oxide catalysts, Ind. Eng. Chem. Res. 51 (2012)
 1029 11828–11840. doi:10.1021/ie3007962.

- B. Moghtaderi, Review of the recent chemical looping process developments for novel
 energy and fuel applications, Energy and Fuels. 26 (2012) 15–40. doi:10.1021/ef201303d.
- 1032 [24] N. Saithong, S. Authayanun, Y. Patcharavorachot, A. Arpornwichanop, Thermodynamic
 1033 analysis of the novel chemical looping process for two-grade hydrogen production with CO2
 1034 capture, Energy Convers. Manag. 180 (2019) 325–337.
 1035 doi:10.1016/j.enconman.2018.11.003.
- [25] C. Agrafiotis, M. Roeb, C. Sattler, A review on solar thermal syngas production via redox pair-based water/carbon dioxide splitting thermochemical cycles, Renew. Sustain. Energy Rev. 42 (2015) 254–285. doi:10.1016/j.rser.2014.09.039.
- 1039 [26] D. Yadav, R. Banerjee, A review of solar thermochemical processes, Renew. Sustain. Energy
 1040 Rev. 54 (2016) 497–532. doi:10.1016/j.rser.2015.10.026.
- 1041 [27] P.T. Krenzke, J.R. Fosheim, J.H. Davidson, Solar fuels via chemical-looping reforming, Sol.
 1042 Energy. 156 (2017) 48–72. doi:10.1016/j.solener.2017.05.095.
- 1043 [28] S. Chuayboon, S. Abanades, S. Rodat, Syngas production via solar-driven chemical looping 1044 methane reforming from redox cycling of ceria porous foam in a volumetric solar reactor, 1045 Chem. Eng. J. 356 (2018) 756–770. doi:10.1016/j.cej.2018.09.072.
- 1046 [29] A.E. Farooqui, A.M. Pica, P. Marocco, D. Ferrero, A. Lanzini, S. Fiorilli, J. Llorca, M.
 1047 Santarelli, Assessment of kinetic model for ceria oxidation for chemical-looping CO2
 1048 dissociation, Chem. Eng. J. 346 (2018) 171–181. doi:10.1016/j.cej.2018.04.041.
- [30] F. Liu, Cerium oxide promoted oxygen carrier development and scale modeling study for
 chemical looping comustion, University of Kentucky, 2013.
 https://uknowledge.uky.edu/cgi/viewcontent.cgi?article=1029&context=me_etds.
- 1052 [31] K.J. Warren, J.R. Scheffe, Kinetic insights into the reduction of ceria facilitated via the
 partial oxidation of methane, Mater. Today Energy. 9 (2018) 39–48.
 doi:10.1016/j.mtener.2018.05.001.
- J. Kim, C.A. Henao, T.A. Johnson, D.E. Dedrick, J.E. Miller, E.B. Stechel, C.T. Maravelias,
 Methanol production from CO2 using solar-thermal energy: process development and
 techno-economic analysis, Energy Environ. Sci. 4 (2011) 3122. doi:10.1039/c1ee01311d.
- IO58 [33] J. Kim, T.A. Johnson, J.E. Miller, E.B. Stechel, C.T. Maravelias, Fuel production from CO2 using solar-thermal energy: system level analysis, Energy Environ. Sci. 5 (2012) 8417.
 IO60 doi:10.1039/c2ee21798h.
- [34] C. Falter, V. Batteiger, A. Sizmann, Climate Impact and Economic Feasibility of Solar
 Thermochemical Jet Fuel Production, Environ. Sci. Technol. 50 (2016) 470–477.
 doi:10.1021/acs.est.5b03515.
- [35] T.C. Davenport, C.K. Yang, C.J. Kucharczyk, M.J. Ignatowich, S.M. Haile, Maximizing fuel
 production rates in isothermal solar thermochemical fuel production, Appl. Energy. 183
 (2016) 1098–1111. doi:10.1016/j.apenergy.2016.09.012.
- 1067 [36] C. Falter, R. Pitz-Paal, Energy analysis of solar thermochemical fuel production pathway
 1068 with a focus on waste heat recuperation and vacuum generation, Sol. Energy. 176 (2018)
 1069 230–240. doi:10.1016/J.SOLENER.2018.10.042.
- 1070 [37] S.H. Park, C.S. Lee, Applicability of dimethyl ether (DME) in a compression ignition engine
 1071 as an alternative fuel, Energy Convers. Manag. 86 (2014) 848–863.
 1072 doi:10.1016/j.enconman.2014.06.051.

- 1073 [38] E. Catizzone, G. Bonura, M. Migliori, F. Frusteri, G. Giordano, CO2 recycling to dimethyl
 1074 ether: State-of-the-art and perspectives, Molecules. 23 (2018) 1–28.
 1075 doi:10.3390/molecules23010031.
- 1076 [39] W. Ying, L. Genbao, Z. Wei, Z. Longbao, Study on the application of DME/diesel blends in a diesel engine, Fuel Process. Technol. 89 (2008) 1272–1280.
 1078 doi:10.1016/j.fuproc.2008.05.023.
- 1079 [40] A. Bakhtyari, M.R. Rahimpour, Methanol to Dimethyl Ether, in: A. Basile, F. Dalena (Eds.),
 1080 Methanol Sci. Eng., Elsevier B.V., 2017: pp. 281–311. doi:10.1016/B978-0-444-63903 1081 5.00010-8.
- 1082 [41] T. Ogawa, N. Inoue, T. Shikada, Y. Ohno, Direct Dimethyl Ether Synthesis, J. Nat. Gas
 1083 Chem. 12 (2003) 219–227.
- T.A. Semelsberger, R.L. Borup, H.L. Greene, Dimethyl ether (DME) as an alternative fuel, J.
 Power Sources. 156 (2006) 497–511. doi:10.1016/j.jpowsour.2005.05.082.
- [43] Department of Energy and Climate Change, DECC Fossil Fuel Price Projections, London,
 2013.
- 1088 [44] W.B. Group, Commodity Markets Outlook, Washington, DC, 2018.
 1089 doi:10.1017/CBO9781107415324.004.
- [45] G. M, K. R, J. E, Production of methanol and dimethyl ether from biomass derived syngas: a comparison of the different synthesis pathways by means of flowsheet simulation, in: 23rd
 Eur. Symp. Comput. Aided Process Eng. (ESCAPE 23) 9-12 June, 2013, 2013: p. 2013.
 doi:10.1016/B978-0-7506-3560-8.50001-9.
- [46] F. Ren, J.-F. Wang, H.-S. Li, Direct mass production technique of dimethyl ether from
 synthesis gas in a circulating slurry bed reactor, Stud. Surf. Sci. Catal. 159 (2006) 489–492.
 https://www.scopus.com/inward/record.uri?eid=2-s2.033745792734&partnerID=40&md5=25145520ec4d439b01fe6100b67846b8.
- [47] T.A. Adams, J.H. Ghouse, Polygeneration of fuels and chemicals, Curr. Opin. Chem. Eng. 10
 (2015) 87–93. doi:10.1016/j.coche.2015.09.006.
- [48] K. Saravanan, H. Ham, N. Tsubaki, J.W. Bae, Recent progress for direct synthesis of
 dimethyl ether from syngas on the heterogeneous bifunctional hybrid catalysts, Appl. Catal.
 B Environ. 217 (2017) 494–522. doi:10.1016/j.apcatb.2017.05.085.
- [49] A. Hankin, N. Shah, Process exploration and assessment for the production of methanol and dimethyl ether from carbon dioxide and water, Sustain. Energy Fuels. 00 (2017) 1–16.
 doi:10.1039/C7SE00206H.
- [50] K. Li, H. Wang, Y. Wei, Syngas generation from methane using a chemical-looping concept:
 A review of oxygen carriers, J. Chem. (2013). doi:10.1155/2013/294817.
- 1108 [51] D.R. Simbeck, A.D. Karp, R.L. Dickenson, Syngas production for gas-to-liquids applications: Technologies, issues and outlook., 1998.
 1110 https://web.anl.gov/PCS/acsfuel/preprint archive/Files/Merge/Vol-45_1-0003.pdf.
- 1111 [52] S. Luo, L. Zeng, D. Xu, M. Kathe, E. Chung, N. Deshpande, L. Qin, A. Majumder, T.L.
 1112 Hsieh, A. Tong, Z. Sun, L.S. Fan, Shale gas-to-syngas chemical looping process for stable
 1113 shale gas conversion to high purity syngas with a H2:CO ratio of 2:1, Energy Environ. Sci. 7
 1114 (2014) 4104–4117. doi:10.1039/c4ee02892a.

- 1115 [53] A. Lerner, M.J. Brear, J.S. Lacey, R.L. Gordon, P.A. Webley, Life cycle analysis (LCA) of
 1116 low emission methanol and di-methyl ether (DME) derived from natural gas, Fuel. 220
 1117 (2018) 871–878. doi:10.1016/j.fuel.2018.02.066.
- 1118 [54] H.M. Shim, S.J. Lee, Y.D. Yoo, Y.S. Yun, H.T. Kim, Simulation of DME synthesis from coal syngas by kinetics model, Korean J. Chem. Eng. 26 (2009) 641–648.
 1120 doi:10.1007/s11814-009-0107-9.
- [55] G.H. Graaf, J.G.M. Winkelman, Chemical Equilibria in Methanol Synthesis Including the
 Water-Gas Shift Reaction: A Critical Reassessment, Ind. Eng. Chem. Res. 55 (2016) 5854–
 5864. doi:10.1021/acs.iecr.6b00815.
- 1124 [56] M.K. Cohce, I. Dincer, M.A. Rosen, Energy and exergy analyses of a biomass-based
 1125 hydrogen production system, Bioresour. Technol. 102 (2011) 8466–8474.
 1126 doi:10.1016/j.biortech.2011.06.020.
- 1127 [57] A.M. Eltony, H.G. Park, S.X. Wang, J. Kong, I.L. Chuang, Motional heating in a graphene1128 coated ion trap, Nano Lett. 14 (2014) 5712–5716. doi:10.1021/nl502468g.
- 1129 [58] D.-Y. Peng, D.B. Robinson, A New Two-Constant Equation of State, Ind. Eng. Chem.
 1130 Fundam. 15 (1976) 59–64. doi:10.1021/i160057a011.
- 1131 [59] L. Fanxing, Z. Liang, V.L. G., Y. Zachary, F. Liang-Shih, Syngas chemical looping
 1132 gasification process: Bench-scale studies and reactor simulations, AIChE J. 56 (2009) 2186–
 1133 2199. doi:10.1002/aic.12093.
- 1134 [60] I. Barin, Thermochemical data of pure substances, (1995) 1885.
 1135 doi:10.1002/9783527619825.
- 1136 [61] M.B. Berkenpas, J.J. Fry, K. Kietzke, E.S. Rubin, IECM User Documentation : User Manual 1137 IECM User Documentation : User Manual, (2018) 509–515.
- 1138[62]CAESAR project European best practice guidelines for assessment of CO2 capture1139technologies., 2011. https://www.sintef.no/globalassets/project/decarbit/d-1-4-11403_euro_bp_guid_for_ass_co2_cap_tech_280211.pdf.
- 1141 [63] M.N. Khan, T. Shamim, Influence of Specularity Coefficient on the Hydrodynamics and
 1142 Bubble Statistics of an Annular Fluidized Bed Reactor, Energy Procedia. 105 (2017) 1998–
 1143 2003. doi:10.1016/j.egypro.2017.03.573.
- 1144 [64] D.K. Bhunya, Simulation Study of Cryogenic Air Separation Unit Using Aspen Hysys At
 1145 Rourkela Steel Plant Master of Technology in (Cryogenic and Vacuum Technology)
 1146 National Institute of Technology Rourkela, National Institute of Technology Rourkela, 2014.
 1147 http://ethesis.nitrkl.ac.in/5971/1/E-138.pdf.
- I148 [65] J. Aprilia, K. Kolmetz, Air seperation units (Engineering design guideline) KLM Technology
 Group-Practical Engineering Guidelines for Processing Plant Solutions, 2013.
 http://kolmetz.com/pdf/EDG/ENGINEERING_DESIGN_GUIDELINE_Air_Seperation_Uni
 ts_Rev01web.pdf.
- 1152 [66] V. Raibhole, S. Sapali, Simulation of Medium Purity Gaseous Oxygen Cryogenic Plant for
 1153 Biomass Gasification by Aspen Plus, Open Access Sci. Reports. 1:343 (2012).
 1154 doi:10.4172/scientificreports.343.
- 1155 [67] B.J.P. Buhre, L.K. Elliott, C.D. Sheng, R.P. Gupta, T.F. Wall, Oxy-fuel combustion
 1156 technology for coal-fired power generation, Prog. Energy Combust. Sci. 31 (2005) 283–307.
 1157 doi:10.1016/j.pecs.2005.07.001.

- 1158 [68] D. Cocco, A. Pettinau, G. Cau, Energy and economic assessment of IGCC power plants
 1159 integrated with DME synthesis processes, Proc. Inst. Mech. Eng. Part A J. Power Energy.
 1160 220 (2006) 95–102. doi:10.1243/095765006X76027.
- 1161 [69] S. Mukherjee, P. Kumar, A. Yang, P. Fennell, Energy and exergy analysis of chemical
 1162 looping combustion technology and comparison with pre-combustion and oxy-fuel
 1163 combustion technologies for CO2 capture, J. Environ. Chem. Eng. 3 (2015) 2104–2114.
 1164 doi:10.1016/j.jece.2015.07.018.
- 1165 [70] Y. Khojasteh-Salkuyeh, New Polygeneration Processes for Power Generation and Liquid
 1166 Fuel Production with Zero CO2 Emissions, McMaster University, 2015.
- 1167 [71] M. Pozzo, A. Lanzini, M. Santarelli, Enhanced biomass-to-liquid (BTL) conversion process
 1168 through high temperature co-electrolysis in a solid oxide electrolysis cell (SOEC), Fuel. 145
 1169 (2015) 39–49. doi:10.1016/j.fuel.2014.12.066.
- 1170 [72] K.L. Ng, D. Chadwick, B.A. Toseland, Kinetics and modelling of dimethyl ether synthesis
 1171 from synthesis gas, Chem. Eng. Sci. 54 (1999) 3587–3592. doi:10.1016/S00091172 2509(98)00514-4.
- 1173 [73] F. Dadgar, R. Myrstad, P. Pfeifer, A. Holmen, H.J. Venvik, Direct dimethyl ether synthesis
 1174 from synthesis gas: The influence of methanol dehydration on methanol synthesis reaction,
 1175 Catal. Today. 270 (2016) 76–84. doi:10.1016/j.cattod.2015.09.024.
- 1176 [74] C. Arcoumanis, C. Bae, R. Crookes, E. Kinoshita, The potential of di-methyl ether (DME) as
 an alternative fuel for compression-ignition engines: A review, Fuel. 87 (2008) 1014–1030.
 doi:10.1016/j.fuel.2007.06.007.
- IT79 [75] J. Sun, G. Yang, Y. Yoneyama, N. Tsubaki, Catalysis chemistry of dimethyl ether synthesis,
 ACS Catal. 4 (2014) 3346–3356. doi:10.1021/cs500967j.
- 1181 [76] R.S. Treptow, Le Chatelier's Principle, J. Chem. Educ. 57 (1980) 417–419.
 1182 doi:10.1021/ed057p417.
- 1183 [77] F. Pontzen, W. Liebner, V. Gronemann, M. Rothaemel, B. Ahlers, CO2-based methanol and
 1184 DME Efficient technologies for industrial scale production, Catal. Today. 171 (2011) 242–
 1185 250. doi:10.1016/j.cattod.2011.04.049.
- 1186 [78] M. Tobiszewski, M. Marć, A. Gałuszka, J. Namiešnik, Green chemistry metrics with special reference to green analytical chemistry, Molecules. 20 (2015) 10928–10946. doi:10.3390/molecules200610928.
- [79] A. Hankin, N. Shah, Process exploration and assessment for the production of methanol and dimethyl ether from carbon dioxide and water, in: A. Basile, F. Dalena (Eds.), Methanol Sci.
 Eng., Elsevier B.V., 2017: pp. 1541–1556. doi:10.1039/C7SE00206H.
- [80] M.C. Gutiérrez, J.M. Rosas, M.A. Rodríguez-Cano, I. López-Luque, J. Rodríguez-Mirasol,
 T. Cordero, Strategic situation, design and simulation of a biorefinery in Andalusia, Energy
 Convers. Manag. 182 (2019) 201–214. doi:10.1016/j.enconman.2018.12.038.
- 1195[81]K.M. Vanden Bussche, G.F. Froment, A Steady-State Kinetic Model for Methanol Synthesis1196and the Water Gas Shift Reaction on a Commercial Cu/ZnO/Al2O3Catalyst, J. Catal. 1611197(1996) 1–10. doi:10.1006/jcat.1996.0156.
- 1198 [82] G. Berčič, J. Levec, Catalytic Dehydration of Methanol to Dimethyl Ether. Kinetic
 1199 Investigation and Reactor Simulation, Ind. Eng. Chem. Res. 32 (1993) 2478–2484.
 1200 doi:10.1021/ie00023a006.

R.S. Schiffino, R.P. Merrill, A mechanistic study of the methanol dehydration reaction on 1201 [83] .gamma.-alumina catalyst, J. Phys. Chem. 97 (1993) 6425-6435. doi:10.1021/j100126a017. 1202 G.W. K. Klier, Simmons, Catalytic Synthesis of Methanol from CO / H2, J. Catal. 360 1203 [84] (1982) 343-360. doi:https://doi.org/10.1016/0021-9517(82)90040-9. 1204 G.R. Moradi, F. Yaripour, P. Vale-Sheyda, Catalytic dehydration of methanol to dimethyl 1205 [85] ether over mordenite catalysts, Fuel Process. Technol. 91 (2010) 461-468. 1206 doi:10.1016/j.fuproc.2009.12.005. 1207 [86] O. Y, S. T, O. T, O. M, M. M, F. K, New Clean Fuel from Coal-Dimethyl Ether, in: 213th 1208 ACS Natl. Meet. San Fr., 1997: pp. 1–5. 1209 D.Y.C. Leung, G. Caramanna, M.M. Maroto-Valer, An overview of current status of carbon 1210 [87] 1211 dioxide capture and storage technologies, Renew. Sustain. Energy Rev. 39 (2014) 426-443. doi:10.1016/j.rser.2014.07.093. 1212 A.D. Curzons, D.J.C. Constable, D.N. Mortimer, V.L. Cunningham, So you think your 1213 [88] process is green, how do you know? - Using principles of sustainability to determine what is 1214 1215 green - A corporate perspective, Green Chem. 3 (2001) 1-6. doi:10.1039/b007871i. [89] R.A. Sheldon, Metrics of Green Chemistry and Sustainability: Past, Present, and Future, 1216 ACS Sustain. Chem. Eng. 6 (2018) 32-48. doi:10.1021/acssuschemeng.7b03505. 1217 [90] I.C. Kemp, Pinch analysis and process integration: A user guide on process integration for 1218 the efficient use of energy, Pinch Anal. Process Integr. (2007) 416. 1219 doi:http://dx.doi.org/10.1016/B978-075068260-2.50003-1. 1220 T.J. Kotas, Chapter 2 - Basic exergy concepts BT - The Exergy Method of Thermal Plant [91] 1221 Analysis, in: Butterworth-Heinemann, 1985: pp. 29-56. doi:https://doi.org/10.1016/B978-0-1222 408-01350-5.50009-X. 1223 A.P. Hinderink, F.P.J.M. Kerkhof, A.B.K. Lie, J. De Swaan Arons, H.J. Van Der Kooi, 1224 [92] Exergy analysis with a flowsheeting simulator - I. Theory; calculating exergies of material 1225 1226 streams, Chem. Eng. Sci. 51 (1996) 4693-4700. doi:10.1016/0009-2509(96)00220-5. J.Y. Xiang, M. Cal, M. Santarelli, Calculation for physical and chemical exergy of flows in 1227 [93] systems elaborating mixed-phase flows and a case study in an IRSOFC plant, 115 (2004) 1228 101-115. doi:10.1002/er.953. 1229 J. Szargut, Chemical Exergies of the Elements, Appl. Energy. 32 (1989) 269–286. 1230 [94] Kristen Gerdes, John Haslbeck, Norma Kuehn, Eric Lewis, Lora L. Pinkerton, Mark Woods, 1231 [95] James Simpson, Marc J. Turner, Elsy Varghese, Cost and Performance Baseline for Fossil 1232 Energy Plants Volume 1: Bituminous Coal and Natural Gas to Electricity, Doe/Netl-1233 2010/1397. 1 (2010). doi:DOE/NETL-2010/1397. 1234 K. Gerdes, W.M. Summers, J. Wimer, Cost Estimation Methodology for NETL Assessments 1235 [96] of Power Plant Performance, Doe/Netl-2011/1455. (2011) 26. http://www.netl.doe.gov/File 1236 Library/research/energy analysis/publications/QGESSNETLCostEstMethod.pdf. 1237 J.A.S. Richard Turton, Richard C. Bailie, Wallace B. Whiting, Analysis, Synthesis and 1238 [97] Design of Chemical Processes Third Edition, 2013. doi:10.1017/CBO9781107415324.004. 1239 [98] Politecnico di Milano - CAESER Project, Enabling advanced pre-combustion capture 1240 1241 techniques and plants; in deliverable D.1.4.3: European best practice guidelines for 1242 assessment of CO2 capture technologies, 2011. 41

- 1243 https://www.sintef.no/globalassets/project/decarbit/d-1-4-
- 1244 3_euro_bp_guid_for_ass_co2_cap_tech_280211.pdf.
- [99] M. Yang, International Energy Agency Working Paper Series Modeling Investment Risks
 and Uncertainties with Real Options Approach Modeling Investment Risks and Uncertainties
 with Real Options Approach, Structure. 13 (2007) 1120–37.
- 1248 http://www.iea.org/Textbase/publications/free_new_Desc.asp?PUBS_ID=1857.
- [100] F. Watson, EU CO2 price to hit Eur30/mt by end 2019: Bank of America, Spglobal.Com.
 (2018). https://www.spglobal.com/platts/en/market-insights/videos/market-moverseurope/112618-oil-price-decline-takes-center-stage (accessed December 4, 2018).
- [101] World Bank Group, State and Trends of Carbon Pricing 2018, Washington, DC, 2018.
 doi:10.1596/978-1-4648-1292-7.
- [102] S. Ambrogi, EU carbon prices could double by 2021 and quadruple by 2030,
 Https://Www.Carbontracker.Org. (2018). https://www.carbontracker.org/eu-carbon-prices could-double-by-2021-and-quadruple-by-2030/ (accessed December 4, 2018).
- [103] Z. Said, A.A. Alshehhi, A. Mehmood, Predictions of UAE's renewable energy mix in 2030,
 Renew. Energy. 118 (2018) 779–789. doi:10.1016/j.renene.2017.11.075.
- [104] J. Bordoff, N. Kaufman, A Federal US Carbon Tax: Major Design Decisions and Implications, Joule. 2 (2018) 2487–2491. doi:10.1016/j.joule.2018.11.020.
- [105] K.C. de Bruin, A.M. Yakut, The Economic and Environmental Impacts of Increasingarbon tax the Irish Carbon Tax: The Economic and Social Research Institute., 2018. doi:https://doi.org/10.26504/rs79.
- [106] B. Dolter, N. Rivers, The cost of decarbonizing the Canadian electricity system, Energy
 Policy. 113 (2018) 135–148. doi:10.1016/j.enpol.2017.10.040.
- [107] The Full Costs of Electricity Provision; NEA No. 7298: Nuclear Energy Agency,
 Organisation for economic co-operation and development, 2018. http://www.oecd nea.org/ndd/pubs/2018/7298-full-costs-2018.pdf.
- [108] A. Mezősi, L. Szabó, S. Szabó, Cost-efficiency benchmarking of European renewable
 electricity support schemes, Renew. Sustain. Energy Rev. 98 (2018) 217–226.
 doi:10.1016/j.rser.2018.09.001.
- [109] M. Johannes, T. Jessica, H. Niklas, S. Charlotte, P. Simon, N. Sebastian, L. Simon, S. Noha,
 S. Thomas, K. Christoph, Levelized Cost of Electricity Renewable Energy Technologies:
 Fraunhofer ISE, 2013. www.ise.fraunhofer.de.
- [110] N. Gal, I. Milstein, A. Tishler, C.K. Woo, Investment in electricity capacity under fuel cost
 uncertainty: Dual-fuel and a mix of single-fuel technologies, Energy Policy. 126 (2019) 518–
 532. doi:10.1016/j.enpol.2018.10.040.
- 1278 [111] Executive summary: Projected costs of generating electricity, 2015. doi:10.1007/s00247-002 1279 0666-y.
- [112] P.N. Dyer, R.E. Richards, S.L. Russek, D.M. Taylor, Ion transport membrane technology for
 oxygen separation and syngas production, Solid State Ionics. 134 (2000) 21–33.
 doi:10.1016/S0167-2738(00)00710-4.
- [113] R. Dindorf, Estimating potential energy savings in compressed air systems, Procedia Eng. 39 (2012) 204–211. doi:10.1016/j.proeng.2012.07.026.