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A cradle-to-gate life cycle assessment of green methanol production using direct air capture†

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This study presents a comprehensive cradle-to-gate life cycle assessment (LCA) of synthetic methanol production, integrating low-temperature solid sorbent direct air capture (DAC) systems with renewable energy sources and green hydrogen to evaluate the environmental impacts of various renewable energy configurations for powering the DAC-to-methanol synthesis processes. Renewable energy-powered configurations result in significantly lower greenhouse gas (GHG) emissions than traditional methanol production methods and DAC systems powered by conventional grid energy. Energy configurations analyzed are current US grid mix, solar photovoltaic (PV) in Alabama and Arizona, USA, onshore wind, run-of-river hydroelectric, and geothermal. Notably, hydroelectric and wind power in the western United States emerge as the most sustainable options, showing the lowest global warming potential (GWP) impacts at -2.53 and -2.39 kg CO₂ eq. per kg methanol produced, respectively, in contrast to the $+0.944$ kg CO₂ eq. from traditional steam methane reforming. Furthermore, this research investigates the use of various heat sources for regenerating low-temperature solid sorbent DAC, emphasizing the potential integration of new experimental results of novel microwave-based regeneration compared to industrial waste heat. Through the analysis of renewable energy scenarios and DAC regeneration heat sources, the research emphasizes the pivotal role of sustainable energy sources in climate change mitigation. This study introduces a new approach by comparing both various renewable energy sources and DAC heat sources to identify the most optimal configurations. This work is also distinguished by its integration of new experimental data on microwave DAC regeneration, offering a unique contribution to the existing body of knowledge. This LCA scrutinizes the environmental impacts of renewably powered DAC-to-methanol systems and compares them with traditional methanol production methods, revealing the significant potential for carbon neutrality. The findings highlight the importance of strategic technology and energy source optimization to minimize environmental impacts, thus guiding the scaling up of DAC and renewable energy technologies for effective climate mitigation. By recognizing the environmental advantages of integrating renewable energy sources with DAC-to-methanol technologies, this research marks a significant step forward in advancing DAC technology and pushes the boundaries of green methanol production toward true sustainability.

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Introduction

The urgency to address climate change has escalated with rising global carbon dioxide (CO₂) levels, pushing Earth's climate system towards a critical state. The Intergovernmental Panel on Climate Change (IPCC) highlights the impacts of human-induced climate change on global weather patterns, including

droughts, heatwaves, and extreme precipitation, attributing these changes to the significant increase in measured global temperatures driven by human activity.^{1,2} Despite international efforts to curb emissions, traditional mitigation strategies are falling short of the ambitious targets set by the Paris agreement, while fossil fuels continue to demonstrate an importance to global economies. To bridge this gap, synthetic fuel production is a promising area of research, particularly when integrated with net-negative CO₂ technologies such as direct air capture.³ DAC, when considered with its full life cycle emissions, holds the potential to offset current and past emissions, emerging as a crucial technology in the quest for net-zero emissions.⁴ This technology is poised to become as significant

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as solar photovoltaics, wind energy, batteries, and electrolyzers in the arsenal of climate change mitigation tools.⁵ Additionally, CO₂ which is removed from ambient air through DAC can be used as feedstock to create synthetic fuel, which alleviates the burden of introducing new CO₂ to the carbon cycle through traditional fossil fuels.

Exploring methanol synthesis through renewable energy and advanced carbon capture technologies highlights a strategic approach in reducing GHG emissions and fostering sustainable chemical production. Nemet and Brandt⁶ along with Taghdisian *et al.*⁷ study the economic and environmental impacts of DAC and its eco-friendly design within methanol synthesis frameworks. These considerations are furthered by Daggash *et al.*,⁸ who emphasize the climate mitigation effectiveness of DAC through a comparative analysis of power-to-fuel and power-to-DAC strategies. In parallel, the environmental advantages of integrating clean energy into such processes are explored by the LCAs conducted by Chen *et al.*⁹ and Al-Qahtani *et al.*,¹⁰ while Meunier *et al.*¹¹ and Khojasteh-Salkuyeh *et al.*¹² emphasize the significance of CO₂ valorization and renewable energy advancements. Sollai *et al.*¹³ and Cormos¹⁴ further assess the techno-economic viability, illustrating how policy and technological progress are making green methanol increasingly competitive.

For specific production methodologies, Lin *et al.*¹⁵ assess methanol production through various CO₂ reduction approaches, pinpointing the substantial environmental impact of electrochemical routes due to their high energy demands. Yet, they advocate for the mitigation of these impacts through renewable energy integration. This perspective is echoed in the analyses conducted by Rigamonti and Brivio,¹⁶ Luu *et al.*,¹⁷ Schreiber *et al.*,¹⁸ Zang *et al.*,¹⁹ Trudewind *et al.*,²⁰ and Ling *et al.*,²¹ who explore the environmental impacts of methanol production using carbon capture and utilization (CCU) technologies and highlight the potential of such processes to significantly reduce GHG emissions, especially when leveraging renewable energy sources and innovative production methods. Nguyen *et al.*²² explore methanol production combining dry methane reforming and partial oxidation to minimize CO₂ emissions. By integrating process design, simulation, and cradle-to-gate LCA, they identify optimal conditions that notably achieve low process emissions at 0.81 kg CO₂ per kg of methanol produced. Khojasteh-Salkuyeh *et al.*²³ demonstrate in their LCA that methanol synthesis from CO₂ through CCU and tri-reforming of methane significantly reduces life cycle GHG emissions to 1.75 and 0.41 kg CO₂ per kg MeOH, respectively, compared to conventional natural gas-based methods. Ryoo *et al.*²⁴ study the CO₂ to methanol conversion processes, evaluating four different methods: coal gasification, coal coking, hydrogenation, and photocatalytic conversion, across various technological readiness levels and find photocatalytic conversion to be the least burdensome for global warming potential (GWP). Photocatalytic conversion is also explored in a LCA by Robbins *et al.*²⁵ Barati *et al.*²⁶ highlight the electrified combined reforming of methane for CO₂ to methanol conversion, showing enhanced efficiency and significant reductions in GHG emissions across Canadian provinces.

The integration of renewable energy within chemical production is further showcased by Biernacki *et al.*,²⁷ who analyze the environmental impacts of converting renewable electricity into methanol. Artz *et al.*²⁸ provide a comprehensive review of CCU technology development, emphasizing the essential role of LCA in optimizing these technologies for economic and environmental sustainability. This narrative is extended by studies like those of Cordero-Lanzac *et al.*^{29,30} and Galusnyak *et al.*,³¹ and Sternberg *et al.*³² who examine the production of green methanol from CO₂ and renewable hydrogen, underscoring the importance of process design and the utilization of renewable energy sources.

Including broader environmental sustainability assessments, Garcia-Garcia *et al.*³³ and Rosental *et al.*³⁴ review the impacts of CCU technologies, including methanol production, advocating for the reduction of global warming impacts through harmonized LCA methodologies. This is complemented by the work of Matzen and Demirel,³⁵ Maimaiti *et al.*,³⁶ and Nizami *et al.*,³⁷ who highlight the environmental benefits of renewable energy-powered methanol production, from wind-derived hydrogen and photovoltaics to solar energy, emphasizing the sector's transition towards sustainability.

Within the context of DAC technology, Van der Giesen *et al.*,³⁸ Liu *et al.*,³⁹ Deutz and Bardow,⁴⁰ Terlouw *et al.*,⁴¹ and Micheli *et al.*⁴² recognize the pivotal role of renewable energy in enhancing the environmental and economic outcomes of DAC technologies. They explore various applications, from liquid hydrocarbon production to aviation sector decarbonization, showcasing the significant benefits of renewable-powered DAC pathways. This collective body of research underlines the necessity of transitioning towards sustainable practices in methanol production, with renewable energy and carbon capture positioned as key components in global climate mitigation efforts, despite the challenges posed by data availability for comprehensive LCA of DAC-based systems. Recent studies highlight the varying efficiencies and social costs associated with different carbon capture technologies. For instance, Jacobson⁴³ emphasizes that synthetic direct air carbon capture and storage and CCU technologies, when powered by natural gas, capture a net of only 10.5–10.8% of CO₂ emissions over 20 years and 20–31% over 100 years due to uncaptured emissions from natural gas combustion and upstream processes.

Building upon the insights of these prior studies, this research marks a distinctive contribution to the field by conducting a comparative analysis of different electrical energy sources and DAC heat sources within the context of synthetic methanol production, including the incorporation of new experimental data on microwave DAC regeneration. This side-by-side examination of these energy sources is novel, enhancing the granularity of LCA methodologies applied to DAC systems and providing a comprehensive assessment of their environmental performance. By integrating new data and contrasting multiple energy pathways, the study sets a new precedent for evaluating the potential of DAC-to-methanol systems, offering unique insights that could shape future research directions,

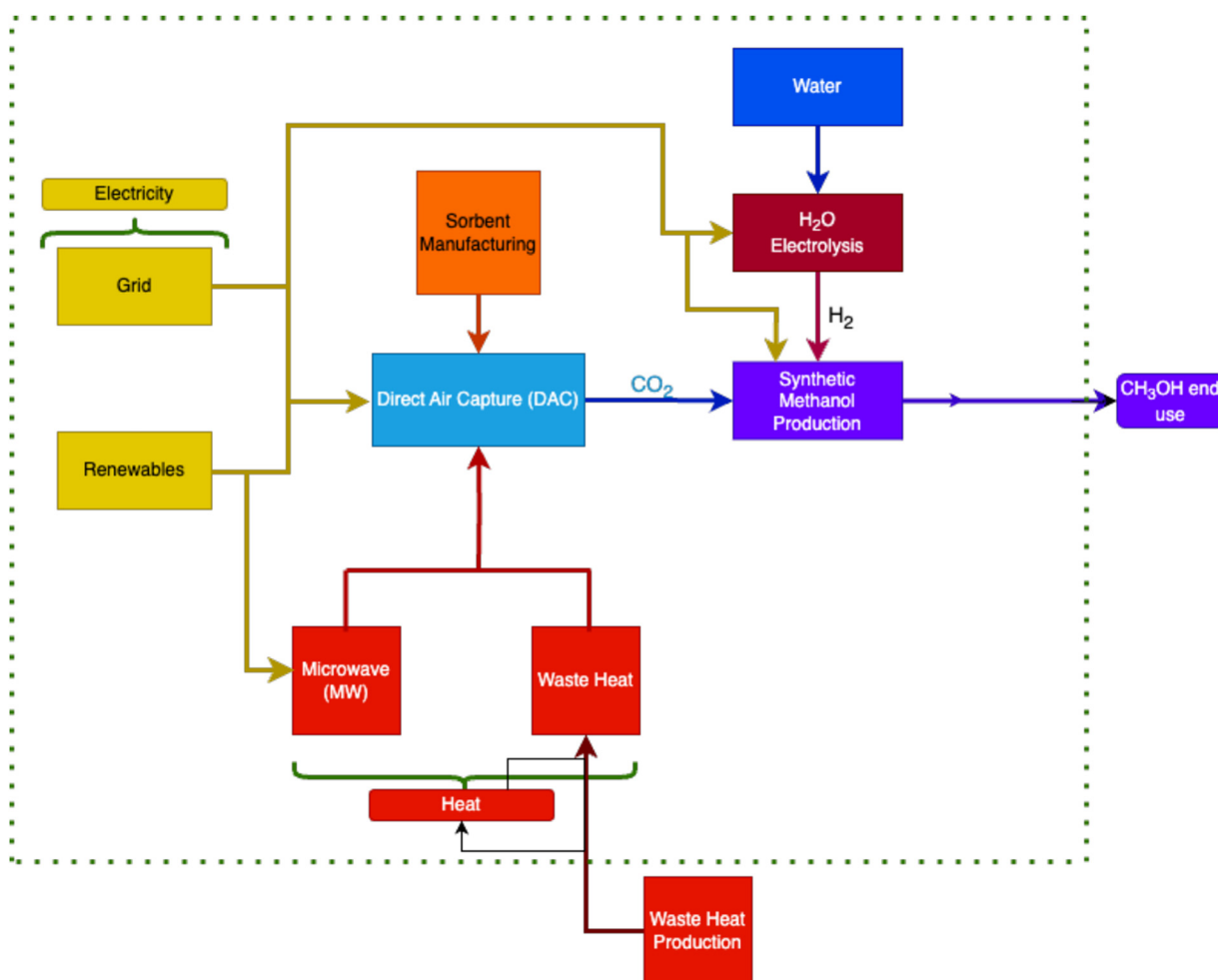


electrolysis processes for H₂ production. Through this study, a comprehensive investigation on the environmental impact of integrating renewably powered DAC with synthetic methanol production is conducted to lower demand for naturally drilled methanol and thereby close the carbon loop. Using captured CO₂ from the DAC plant as an input to synthetic methanol has the added benefit of potentially offsetting the production cost through the sale of green fuel, which could lower the net cost of DAC. This study extends the scope of previous life cycle assessments of DAC systems by focusing on the environmental implications and potential for integrating renewably powered DAC with hydrogen and green methanol production systems.

This study's scope is cradle-to-gate, examining all processes from the extraction of raw materials to production of synthetic methanol. While the end use of methanol is not considered in the scope, all other end of life (EOL) processes associated with DAC and the construction are included. The results of this study will be compared to traditional methanol production methods, also cradle-to-gate, to understand the improvements to the environmental impacts which could be made through

The goal of this work is to analyze multiple sustainable energy scenarios where renewably powered green methanol production and DAC to identify the most sustainable operational methods for GHG reduction. While some sources such as geothermal or hydroelectric would require co-location at a pre-existing or planned plant, others such as PV or wind-powered production could be planned as an off-the-grid, sustainable solution to mitigate climate change in many different areas, from urban to remote. This flexibility is a key advantage to certain renewables which could benefit these processes greatly.

Fig. 1 illustrates the proposed system which contains three main components: the CO₂ hydrolysis methanol production plant, the low temperature solid sorbent DAC plant, and the



Energy Adv., 2024, 3, 2311-2327 | 2313

consumption, construction, process chemical production, waste management, and decommissioning. The assumptions for the construction and EOL treatment of the infrastructure is available in the electronic supplementary information (ESI).†

Aspen Plus software was used to simulate the process flow for synthetic methanol production *via* CO₂ hydrogenation using an adiabatic fixed-bed catalytic reactor based on the processes studied by Sollai *et al.*¹³ and Meunier *et al.*,¹¹ *via* eqn (1).^{11,48}

$$\text{CO}_2 + 3\text{H}_2 \leftrightarrow \text{CH}_3\text{OH} + \text{H}_2\text{O} \quad (1)$$

Fig. 2 illustrates the flowsheet of the methanol production process. The CO_2 input to the process is concentrated by the DAC system while the H_2 is formed through water electrolysis on-site as described in Section 3.3. Prior to entering the reactor, these inputs are mixed and compressed to 6.5 MPa, and they pass through the first heat exchanger to increase to the operating temperature of 210 °C.¹³ Following the reaction, the majority of unreacted CO_2 and H_2 , and the produced H_2O is separated from the outlet stream.¹¹ Most of the remaining H_2O is removed in the distillation column, and the remaining CO_2 is returned to the inlet.¹¹ Overall, 281 kg h⁻¹ of H_2O is calculated to be removed during purification, assumed in the LCA to be sent to water treatment.

The resultant methanol stream is found to have a purity of 95.3%. Most of the separated CO_2 and H_2 is recycled back to the inlet of the reactor, but the methanol stream still contains trace amounts of CO_2 and H_2O ¹¹ which are purged and flared to remove them from the circuit.¹³ This purge flow does increase CO_2 emissions slightly, but this contribution is almost

The second phase of the LCA methodology is to gather all inputs and outputs associated with the unit processes of the system boundary before the impact results can be calculated.⁴⁷ In this study, some of the most important unit processes included are methanol production, DAC operations, energy

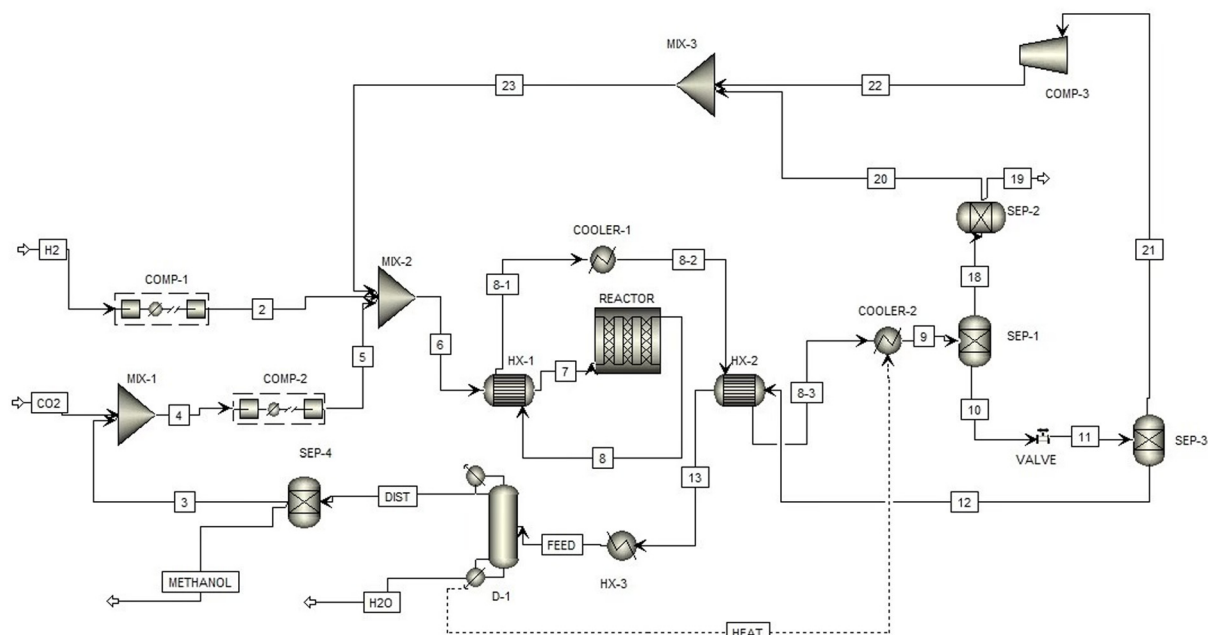


Fig. 2 Process flow for methanol production as considered in this study.

Table 1 Key parameters used in this analysis for methanol production

Parameter	Value
Design capacity	2.63 kt yr ⁻¹
CO ₂ inlet mass flow	723 kg h ⁻¹ (ref. 13)
H ₂ inlet mass flow	104 kg h ⁻¹ (ref. 13)
CO ₂ outlet mass flow to atmosphere	36.32 kg h ⁻¹
H ₂ outlet mass flow to atmosphere	9.64 kg h ⁻¹
H ₂ O outlet mass flow to treatment	281 kg h ⁻¹
Methanol outlet mass flow	500 kg h ⁻¹ (ref. 13)
Electricity load	108.64 kW
Catalyst amount	292.4 kg Cu/Zn/Al ¹³
Catalyst lifetime	4 yr ¹³

negligible since it is only about 3% of the total.¹³ It was also found that there is a CO₂ and H₂ losses to the atmosphere during the separation processes with the flow rates of 36.32 kg h⁻¹ and 9.64 kg h⁻¹, respectively. These atmospheric release purges slightly lower the net amount of CO₂ removed from the air by about 5%, which is accounted for as an air emission.

The catalyst for this reaction is Cu/Zn/Al, which is well documented to be an efficient and economically viable catalyst material.^{11,13,49,50} The selection of this catalyst is based on its proven capability to operate effectively in fixed-bed catalytic reactors at the required temperatures (250–300 °C) and pressures (50–100 bar), conditions that are ideal for methanol synthesis from CO₂ and H₂.¹³ This catalyst has been verified to have a lifetime of at least four years, and eight years has been observed.^{13,51} Four years is assumed for this study.

Table 1 lists the assumptions for the methanol production plant. The inlet mass-flow rates, temperature, operating pressure requirements, and general flows are based on the research by Sollai *et al.*,¹³ while the proportion of unreacted inputs which are recycled derives from Meunier *et al.*¹¹ The resultant CO₂, H₂, and H₂O outlet flows, and electrical load are calculated in Aspen Plus and align well with the previous values as determined by Sollai.¹³ The annual design capacity of the methanol plant is based on the conversion of 4 kt CO₂ per year.

Direct air capture

The type of DAC analyzed is a low-temperature solid sorbent system of an annual capacity of capacity 4000 tons CO₂, similar

to the nominal plant capacity analyzed by Deutz and Bardow.⁵² Auxiliary electrical load for fans and other DAC electronics is assumed to be 0.555 kW h per kg CO₂.⁵³ Fan electricity is well documented to play only a small role in overall energy consumption when compared to regeneration heat.^{40,41,46,53}

The decision to employ DAC with adsorption rather than absorption is primarily driven by the significant advancements and cost efficiencies associated with low-temperature solid sorbents.⁵⁴ Compared to absorption, adsorption processes typically require lower energy inputs for regeneration, particularly when utilizing WH from industrial processes. Recent studies,^{55–57} highlight that the use of solid sorbents can reduce overall heat supply costs, making the technology more economically viable. The continuous improvements in sorbent materials have led to higher CO₂ capture capacities and enhanced stability, further lowering the operational costs and enhancing the commercial feasibility of DAC technologies. The main parameters of the DAC system are included in Table 2.

To compare the life cycle impacts of regeneration heat sources, heat is assumed to be supplied to the DAC system from each of two sources: MW and industrial WH. MW-based heat is based on two the principles of microwave swing desorption (MSD) as studied by Erguvan *et al.*,⁶¹ who examined the potential of using MW technology for the desorption of CO₂ from zeolite 13X. MW as a source for regeneration heat for carbon capture and DAC had previously been proposed by others.^{62–65} Erguvan *et al.* focused on simulated flue gas capture, and subsequent work was performed to determine the regeneration heat requirements of a similar setup but for ambient air through DAC.⁵⁸ During this LCA, MW-based regeneration heat will be a scaled-up DAC regeneration system based on the results of experimental MW DAC work performed by Boylu, Erguvan, and Amini⁵⁸ as shown in Fig. 3.

The efficiency of two key processes in MW-based DAC—adsorption and desorption processes—were investigated using a mono-mode microwave unit. Microwave power and regeneration temperature were varied from 4 to 30 W. Three grams of zeolite 13X were placed in the reactor, and the microwave unit was set to reach a target temperature of 33 °C, 50 °C, and 100 °C.

The sorbent, zeolite 13X, was preconditioned in a conventional oven at 350 °C overnight to eliminate volatiles, following

Table 2 Key parameters used in this analysis for DAC operation

Parameter	Value
DAC design capacity	4 kt CO ₂ per year ⁴⁰
Capacity factor	0.95
DAC equipment lifetime	20 yr ^{40,41,46}
DAC location (PV-powered)	Tuscaloosa, AL, USA and Phoenix, AZ, USA
DAC location (grid-powered)	United States of America
DAC location (wind, geothermal, hydro-powered)	Western United States of America
Base electricity load	0.555 kW h per kg CO ₂ ⁵³
Heat load (MW)	4.8 MJ per kg CO ₂ captured ⁵⁸
Heat load (WH)	7.72 MJ per kg CO ₂ captured ⁵⁹
Electrical source	U.S. grid mix, PV, wind, geothermal, hydroelectric
Heat source	MW, WH
Sorbent type	Zeolite 13X on a cordierite substrate
Sorbent consumption rate	0.788 g per kg CO ₂ captured ⁶⁰

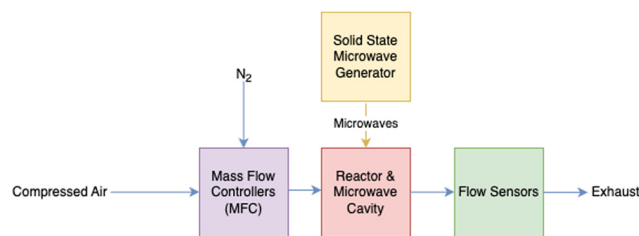


Fig. 3 General flow diagram of the laboratory experimental setup for the DAC with MW regeneration and zeolite 13X sorbent.

the method by Ellison *et al.*⁶⁶ Subsequently, the reactor was purged with nitrogen gas at a flow rate of 200 mL min⁻¹ for 20 minutes to create a CO₂-free environment. During the adsorption phase, compressed air was introduced with a flow rate of 750 mL min⁻¹ to simulate fluidized bed behaviors. The adsorption process was deemed complete when the sorbent reached CO₂ saturation within two hours.

Following adsorption, the feed gas to the reactor was switched to N₂ with the same flow rates to purge any desorbed CO₂. Concurrently, the microwave heating was initiated at each power level to achieve and maintain the desired temperature for regeneration. The power output of the MW generator was adjusted dynamically to maintain the temperature as close as possible to the target. The regeneration phase was concluded once the CO₂ concentration in the effluent gas dropped to 30 ppm. This meticulous control of microwave power ensured that the regeneration temperature remained stable, illustrating the precise management of the desorption process under the influence of microwave heating.

It was found that the most efficient microwave parameters for DAC regeneration are using a MW power setting of 4 W and with a regeneration temperature of 33 °C which achieved a regeneration energy demand of 4.8 MJ per kg CO₂.⁵⁸ This regeneration heat value is assumed for all MW DAC configurations.

Zeolite 13X was selected as the sorbent for the WH case as well to facilitate closer comparison to MW. The details of the inventory for zeolite manufacturing and lifetime are included in the ESI.† When determining the heat requirements for desorption of CO₂ with WH, there is some variance in the literature regarding regeneration heat requirements for zeolite 13X. Generally, sources have demonstrated a heat load of 3–6 MJ per kg CO₂ in temperature ranges of 125–200 °C;^{59,67–70} in this study, we will use the value of 7.72 MJ kg⁻¹ at 100 °C for the case of WH,⁵⁹ as it falls near the middle of the values reported in the literature, and is a temperature of heat readily available in industry.

The availability of WH for DAC desorption is substantial, particularly in industrial settings such as power plants, chemical plants, and refineries. In industrial facilities, including power plants, about 60–70% of the energy produced is discarded as waste heat.⁷¹ This makes WH a practical and cost-effective option for DAC systems. Typically, WH for the proposed DAC system with a capacity of 4 kt per year would require 0.978 MW heat on average. For a 1000 MW fossil fuel

plant at the average efficiency of 36%,⁷¹ this corresponds to up to 640 MW waste heat available. Some previous studies have not considered the burden due to use of WH, considering it free energy.⁴⁰ In this study, an exergy calculation is applied as the allocation to demonstrate the usefulness potential of the steam, as shown in eqn (2). Q is given as the heat requirement of the steam, T_0 is the assumed ambient temperature of 25 °C, and T is the saturated required steam temperature of 100 °C.⁵⁹

WH and MW heating each offer unique benefits for DAC desorption. WH, commonly found in industrial settings, can provide a cost-effective source of thermal energy by repurposing heat that would otherwise be lost. This makes WH a practical option where industrial processes produce ample excess heat. However, its availability and efficiency depend on the specific industrial context and the quality of the heat source, and DAC is limited to a location near an ample WH source. Conversely, MW heating has the potential to offer superior energy efficiency and greater operational flexibility. Even at lab scale, MW heating can achieve lower regeneration energy demands and precise, rapid heating directly to the sorbent material, making it especially useful in locations where WH is unavailable. This technology's ability to deliver targeted heating with lower energy consumption is expected to improve with further research.

Eqn (2) exergy calculation used in the allocation of waste steam, where Q is the heat requirement, T_0 is the ambient temperature, and T is the system temperature, with both temperatures are units of absolute temperature.⁷²

$$Ex = Q \left(1 - \frac{T_0}{T} \right) \quad (2)$$

Hydrogen production

Hydrogen production as an input to the catalytic hydrogenation of CO₂ for synthetic methanol production is assumed to take place on location with methanol production using H₂O electrolysis through a polymer electrolyte membrane (PEM) electrolyzer and using the mass and energy flows as researched by Sollai *et al.*¹³ The selection of this technology was influenced by its quick activation from a cold state, enhanced adaptability, and thus, a more effective integration with variable and intermittent power sources, including renewable energies like solar and wind, which makes this production method easily adaptable to the renewable sources analyzed.¹³ Despite these advantages, PEM electrolysis technology faces challenges, including higher costs and limited operational lifespan. Current systems typically last around 60 000 hours before requiring stack replacement due to degradation, though advancements are expected to extend this to 85 000 hours. The specific energy consumption ranges from 50 to 80 kWh per kilogram of hydrogen produced. For optimal environmental benefits, it is essential to power PEM electrolyzers with renewable energy, as demonstrated in the e-methanol production study by Sollai *et al.*, which utilized overgenerated renewable electricity to minimize carbon footprint and enhance sustainability.¹³

In our study, the produced H₂ is used as an input to the methanol production process. The outlet flow of O₂ is assumed



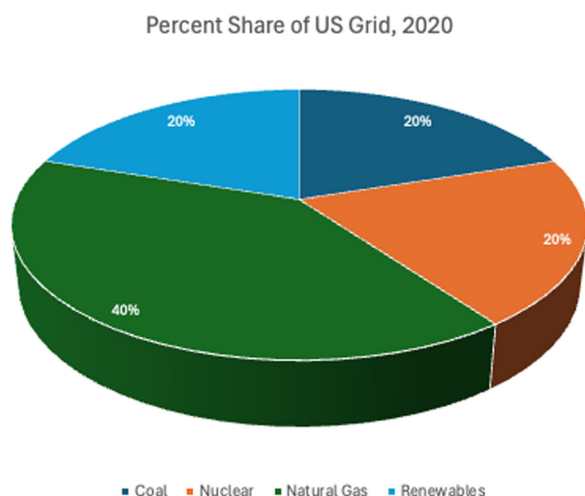
Table 3 Key parameters used in this analysis for hydrogen production

Parameter	Value
Hydrogen production type	H ₂ O PEM electrolysis
H ₂ O inlet mass flow	1037 kg h ⁻¹ (ref. 13)
H ₂ outlet mass flow	104 kg h ⁻¹ (ref. 13)
O ₂ outlet mass flow	821 kg h ⁻¹ (ref. 13)
H ₂ O outlet mass flow to treatment	110 kg h ⁻¹ (ref. 13)
Electricity load	5906 kW ¹³

to be released to the atmosphere as an unwanted product. The construction of the electrolyzer is assumed to be included in the infrastructure construction of the methanol plant. Sensitivity analysis has been conducted to confirm the impacts to the study results based on construction inventories. The mass flow rate of the hydrogen production unit has been listed in Table 3.

Energy scenarios

As discussed earlier, five different electrical energy scenarios, US grid mix, solar photovoltaic, onshore wind power, geothermal power, and run-of-river hydroelectric power, are analyzed in this study. All processes are assumed to take medium voltage electricity, which is standard for most industrial applications. US grid mix is the average energy mix for the United States based on 2020 data.⁷³ By 2020 the US grid has converted power generation by fuel type to about 20% coal, 20% nuclear, 40% natural gas, and 20% renewables as shown in Fig. 4, down from 49% coal, 19% nuclear, 22% natural gas, 2% oil, and just 8% renewables in 2007.⁷⁴ As the United States continues to drive development of renewables, nuclear, and clean natural gas with carbon capture, the grid will continue to improve its carbon footprint. However, the purpose of this research is to identify the most favorable scenarios for implementation of sustainably powered methanol production. Therefore, the US grid is intended to serve mainly as a comparison tool; the results will show that powering these processes through dedicated renewables will have a more favorable impact to the environment.

**Fig. 4** Percent share of US grid production by fuel type, 2020.

For wind, geothermal, and hydroelectric power, ecoinvent 3.8 processes were used.⁷⁵ The assumption is that the methanol and DAC facilities are developed in proximity to pre-existing power plants of the respective type, since the power consumption is relatively small compared to the full capacity of many of these facilities; therefore, construction is only accounted for as a component of the use inventories developed by ecoinvent. Inventories from the Western Electricity Coordinating Council (WECC) for wind, hydroelectric, and geothermal power were used because the WECC reports separately by renewable energy type in ecoinvent while other US sources do not. Wind is assumed to be 1–3 MW onshore turbines which would be more than capable of supplying the electrical load for the facility. According to ecoinvent, as of the year in which data was collected for wind power inventories in 2014, 93.2% of all wind farms in the western United States use 1–3 MW turbines.⁷³ For this study, geothermal power is assumed to be deep well, and hydroelectric is run-of-river type.

PV-based electricity scenarios are set in two locations: Tuscaloosa, Alabama (AL) and Phoenix, Arizona (AZ). Tuscaloosa was chosen since it is a location with medium-high sunlight, medium humidity, and sufficient land area for development. Since the University of Alabama's Decarbonization Laboratory is based in Tuscaloosa, this analysis will help to demonstrate the viability for future renewably-powered DAC projects. Phoenix, Arizona was chosen through an optimization process which analyzed the annual solar irradiation, cloud cover, and available land near utilities infrastructure and support systems to find the most beneficial location for PV in the U.S.^{76,77}

The sizing of PV was completed in both locations for each of the two regeneration heat sources since the total annual electrical load varies depending on heat source. The solar arrays are sized to provide on average the full annual power requirement of all processes: methanol production, hydrogen electrolysis, and DAC. Input parameters and calculated nominal capacities required are shown in Table 4. The solar arrays are assumed to be connected to the grid to supply additional electricity needed for night and times of low PV production while supplying renewable energy to the grid when producing excess for an average annual net grid electrical consumption of zero. All solar irradiation and PV data is provided by the Global Solar Atlas developed by World Bank Group, which uses complex PV modeling algorithms, paired with historical solar

Table 4 Location solar historical data and calculated system sizing, based on data from Global Solar Atlas⁷⁷

Solar Parameter	Tuscaloosa, AL	Phoenix, AZ
Global tilted irradiation	1885 kW h m ⁻² yr ⁻¹	2437 kW h m ⁻² yr ⁻¹
Optimal angle	30°	32°
Azimuth	180°	180°
Power requirement (MW)	38.79 GW h yr ⁻¹	38.79 GW h yr ⁻¹
Power requirement (WH)	33.72 GW h yr ⁻¹	33.72 GW h yr ⁻¹
System capacity (MW)	25 436 kWp	20 491 kWp
System capacity (WH)	22 111 kWp	17 813 kWp
System area (MW)	20 476 m ²	16 495 m ²
System area (WH)	17 799 m ²	14 340 m ²



irradiation data, to estimate monthly and annual electrical energy production based on user inputs.⁷⁷ Construction and EOL processes are fully considered: the construction of the plant includes the ecoinvent infrastructure process of a 570 kWp PV plant scaled based on the nominal PV size required to supply the annual electrical load for the respective energy configuration and reference location in order to capture all the PV electrical infrastructure and construction rather than simply the solar panels. EOL recycling and waste treatment processes are based on research for user-scale smart grid treatment by Rossi *et al.*^{78,79} and Latunussa *et al.*⁸⁰

For all power scenarios, two possibilities for DAC regeneration heat were considered: MW and industrial WH. While each case is considered for purposes of comparison, some are less likely to be implemented in practicality; for instance, if a DAC plant is located nearby a geothermal power plant, it is likely that WH could be used as a heat source as in a current implementation by Climeworks.⁴⁰ Therefore, needing MW-generated heat would likely be less likely than in a remote location powered by PV. These cases are all included in the study for full comparison, but the practicality of implementation depends on the specific application.

Impacts of weather

When exploring the viability of any system powered by renewable energy such as wind or solar, it is imperative to consider the inherent variability and dependency on weather conditions of the primary renewable energy inputs. These sources can fluctuate substantially in output since their availability and efficiency are directly influenced by weather patterns, which can vary significantly over time and location. As a result, the assumptions regarding energy supply for the DAC-to-methanol system are based on average values derived from historical weather and usage data and projected trends. However, this introduces a degree of variability and uncertainty into the energy supply estimates, impacting the overall efficiency and environmental performance of the DAC system. The solar PV cases analyzed have arrays sized to overproduce to the grid during times of high solar irradiation while using grid energy during low production with no net grid usage annually, although in practicality this can vary due to weather. Therefore, when assessing the practicality and reliability of such renewable-powered systems, it is crucial to implement strategies that

mitigate the risks associated with their weather-dependent nature to ensure stable and efficient operations.

The performance of DAC systems using moisture-swing sorbents is notably sensitive to ambient weather conditions, especially temperature and humidity.^{81,82} These systems are more effective in hot, dry climates where the absence of humidity facilitates the release of CO₂ during the regeneration phase. In contrast, operation in cold and moist climates can be less efficient and may require adaptations, such as the addition of a mild thermal swing to improve performance.⁸² Zeolite 13X performance may be significantly degraded in high humidity such as 60% because the interactions between the zeolite and water molecules are so strong that they cannot be removed without significant heating.⁸³ Due to the variable nature of the weather and humidity, actual performance could vary, and careful consideration should be taken when selecting the location for a DAC facility.

Life cycle impact assessment

In this study, a comprehensive LCA has been conducted to evaluate the environmental impacts and efficiency of DAC-to-methanol systems powered by low-carbon energy sources. The inventories for each case were calculated in SimaPro 9.5, and unit processes used were primarily from the ecoinvent 3.8 database.⁷⁵ Based on guidelines from the US DOE regarding conducting a LCA on DAC systems, a modified version of EPA's TRACI version 2.1 method (US, 2008) was used to characterize the impact categories of ozone depletion, global warming, smog formation, acidification, eutrophication, carcinogenics, non-carcinogenics, respiratory effects, ecotoxicity, and fossil fuel depletion.⁴⁷ Since TRACI 2.1 was last updated in 2012, it does not reflect recent GWP characterization factors from the IPCC.⁴⁷ Hence, the TRACI 2.1 method was modified for global warming to add or change the characterizations to reflect the recent IPCC 2021 report.⁸⁴

Table 5 demonstrates the comparative results of methanol production with a MW heat source for each energy scenario, compared to traditional production *via* steam methane reforming (SMR), while Table 6 similarly shows results for the WH cases. Renewably powered MW emerges as the better heat source for environmental impacts, which opens the possibility for more sustainable DAC deployment in remote locations away

Table 5 Life cycle impacts of methanol production from DAC-produced CO₂ with MW regeneration heat and various electrical power sources compared to production from traditional SMR, per kg of synthetic methanol produced

Impact category	Unit	MW, Grid	MW, PV (AL)	MW, PV (AZ)	MW, Wind	MW, Hydro	MW, Geothermal	SMR
Ozone depletion	kg CFC-11 eq.	4.78×10^{-7}	1.27×10^{-7}	1.01×10^{-7}	7.75×10^{-9}	-2.91×10^{-9}	3.74×10^{-8}	2.33×10^{-7}
Global warming	kg CO ₂ eq.	5.57×10^0	-1.36×10^0	-1.60×10^0	-2.39×10^0	-2.53×10^0	-1.47×10^0	9.44×10^{-1}
Smog	kg O ₃ eq.	1.76×10^{-1}	4.26×10^{-2}	2.96×10^{-2}	-1.17×10^{-2}	-1.87×10^{-2}	3.43×10^{-2}	2.51×10^{-2}
Acidification	kg SO ₂ eq.	2.04×10^{-2}	6.50×10^{-3}	5.32×10^{-3}	1.41×10^{-3}	6.98×10^{-4}	4.83×10^{-3}	1.99×10^{-3}
Eutrophication	kg N eq.	4.18×10^{-2}	4.14×10^{-3}	3.21×10^{-3}	1.91×10^{-4}	-5.22×10^{-4}	2.98×10^{-3}	5.51×10^{-4}
Carcinogenics	CTUh	1.48×10^{-6}	1.22×10^{-6}	1.17×10^{-6}	1.16×10^{-6}	9.90×10^{-7}	1.24×10^{-6}	2.48×10^{-8}
Non-carcinogenics	CTUh	3.02×10^{-6}	2.00×10^{-6}	1.86×10^{-6}	1.41×10^{-6}	1.29×10^{-6}	1.54×10^{-6}	5.82×10^{-8}
Respiratory effects	kg PM2.5 eq.	1.43×10^{-2}	2.69×10^{-4}	5.77×10^{-6}	-8.08×10^{-4}	-9.50×10^{-4}	1.63×10^{-4}	2.07×10^{-4}
Ecotoxicity	CTUe	1.89×10^2	1.99×10^2	1.83×10^2	1.48×10^2	1.19×10^2	1.33×10^2	3.65×10^0
Fossil fuel depletion	MJ surplus	7.12×10^0	4.40×10^{-1}	2.41×10^{-1}	-3.73×10^{-1}	-5.29×10^{-1}	1.15×10^{-1}	5.06×10^0

Table 6 Life cycle impacts of methanol production from DAC-produced CO₂ with WH regeneration and various electrical power sources, compared to methanol production from traditional SMR, per kg of synthetic methanol produced

Impact category	Unit	MW, Grid	MW, PV (AL)	MW, PV (AZ)	MW, Wind	MW, Hydro	MW, Geothermal	SMR
Ozone depletion	kg CFC-11 eq.	4.40×10^{-7}	1.35×10^{-7}	1.20×10^{-7}	3.18×10^{-8}	2.25×10^{-8}	5.75×10^{-8}	2.33×10^{-7}
Global warming	kg CO ₂ eq.	4.80×10^0	-1.22×10^0	-1.31×10^0	-2.12×10^0	-2.24×10^0	-1.32×10^0	9.44×10^{-1}
Smog	kg O ₃ eq.	1.58×10^{-1}	4.22×10^{-2}	3.84×10^{-2}	-4.97×10^{-3}	-1.11×10^{-2}	3.50×10^{-2}	2.51×10^{-2}
Acidification	kg SO ₂ eq.	1.86×10^{-2}	6.53×10^{-3}	6.39×10^{-3}	2.11×10^{-3}	1.49×10^{-3}	5.08×10^{-3}	1.99×10^{-3}
Eutrophication	kg N eq.	3.65×10^{-2}	3.78×10^{-3}	3.37×10^{-3}	3.44×10^{-4}	-2.76×10^{-4}	2.76×10^{-3}	5.51×10^{-4}
Carcinogenics	CTUh	1.41×10^{-6}	1.19×10^{-6}	1.20×10^{-6}	1.13×10^{-6}	9.87×10^{-7}	1.21×10^{-6}	2.48×10^{-8}
Non-carcinogenics	CTUh	2.82×10^{-6}	1.93×10^{-6}	1.86×10^{-6}	1.42×10^{-6}	1.32×10^{-6}	1.53×10^{-6}	5.82×10^{-8}
Respiratory effects	kg PM2.5 eq.	1.23×10^{-2}	1.88×10^{-4}	9.33×10^{-5}	-7.48×10^{-4}	-8.72×10^{-4}	9.57×10^{-5}	2.07×10^{-4}
Ecotoxicity	CTUe	1.78×10^2	1.87×10^2	1.75×10^2	1.42×10^2	1.17×10^2	1.29×10^2	3.65×10^0
Fossil fuel depletion	MJ surplus	6.56×10^0	7.47×10^{-1}	6.56×10^{-1}	4.08×10^{-2}	-9.51×10^{-2}	4.65×10^{-1}	5.06×10^0

from WH sources, although co-location with an industrial WH source is shows great improvement in many impact categories over traditional production.

Results in Fig. 5 demonstrate that all renewably powered DAC-to-methanol systems which utilize MW or WH for regeneration are net negative and much more favorable for GWP than traditional methanol production routes. This study finds that the most favorable renewable energy source to power the DAC-to-methanol processes is run-of-river hydroelectric, followed by wind power. These two sources generally were the most favorable among the renewables simply based on the low emissions cataloged in the WECC inventory based on historical plant data.⁷⁵ The PV cases and geothermal power also demonstrate the potential for improvements in GWP impact compared to traditional methanol production *via* SMR.

The difference in impacts between AL and AZ is due to addition to the higher solar irradiation and greater capacity factor associated with Phoenix, which allows smaller sized PV infrastructure to be needed for the required annual capacity. The most favorable DAC regeneration heat scenario is the use of WH from outside industrial processes, as this source provides less burden than the other heat sources. In all cases, the methanol hydrogenation catalyst demonstrated only negligible effects to the overall system life cycle compared to energy usage and construction. Further results for each energy configuration are available in the ESI.[†]

Comparison to traditional methanol production

The results of the study are also compared to traditional methanol production in Tables 5 and 6 through SMR using

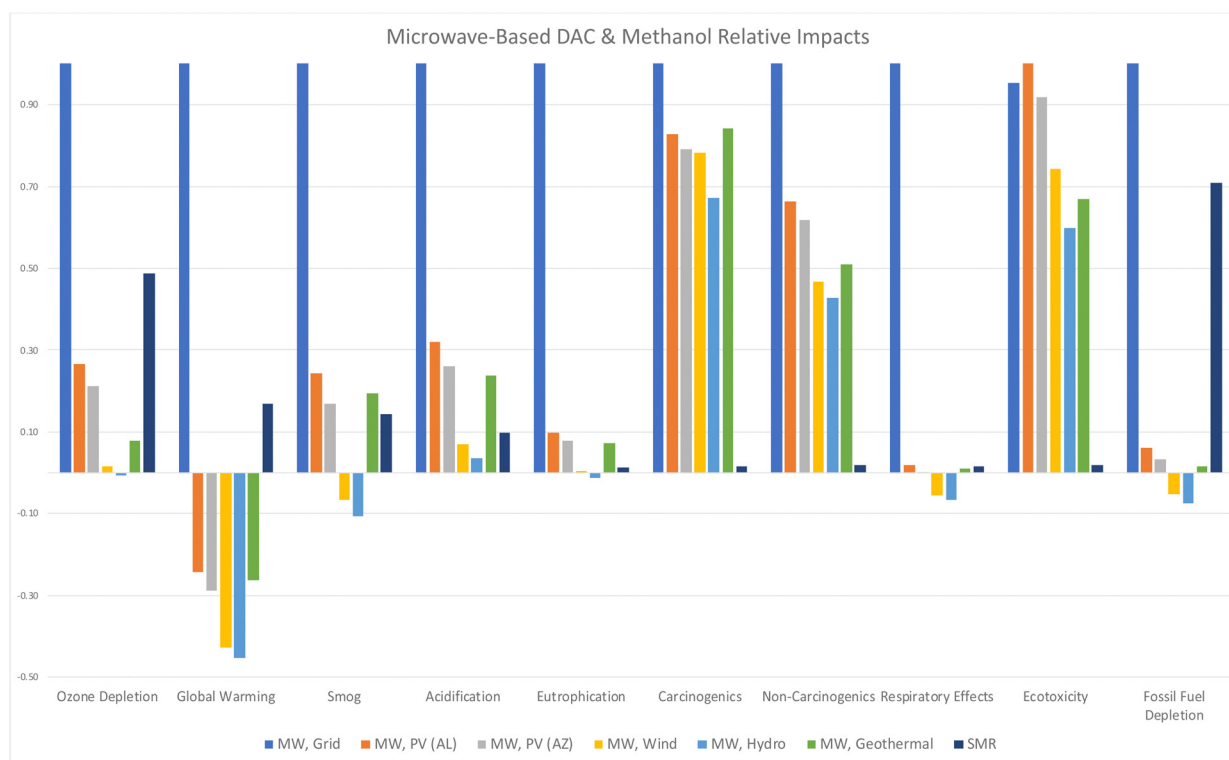


Fig. 5 Relative impacts for methanol production with various energy sources from MW-based DAC compared to traditional production *via* SMR, per kg of methanol produced.



the ecoinvent process for methanol production, which derives its values from literature on plant design and efficiency.⁷⁵ Notably, while all renewably powered cases demonstrate favorable GWP benefits compared to traditional SMR production, SMR shows lower impacts than many of the energy scenarios for smog, acidification, acidification, eutrophication, carcinogenics, non-carcinogenics, and ecotoxicity, as demonstrated in Fig. 5.

The largest contributor to each of these impacts in the DAC-to-methanol system is hydrogen production *via* hydrolysis, due to the high amount of energy required. The electrolysis process for green hydrogen production *via* hydroelectric energy was compared to hydrogen production *via* SMR in Table 7 to identify whether this method could be a favorable replacement process. Results demonstrate that despite the high electrical energy requirement, electrolysis remains less impactful than SMR except for ecotoxicity and carcinogenics. As with the other processes in this DAC-to-methanol system, ensuring electrolysis is integrated with renewable sources is key to minimizing most impacts.

This analysis assumes that the O₂ produced during hydrolysis is compressed and saved as a co-product, which claims a credit based on the process inventory for production *via* cryogenic air separation, the most efficient and cost-effective form of O₂ production.⁷⁵ The total GWP credit for O₂ is substantial at −2.03 kg CO₂ eq. per kg methanol produced. As such, retaining the produced O₂ for industrial use or resale should be considered as some processes may fail to reach net negative emissions, even when powered by renewable energy.

Table 7 Impacts of hydrogen production *via* conventional SMR compared to hydroelectric-powered green hydrogen electrolysis, per kg H₂ produced

Impact category	Unit	Hydrogen <i>via</i> SMR	Electrolysis
Ozone depletion	kg CFC-11 eq.	2.45×10^{-6}	2.28×10^{-8}
Global warming	kg CO ₂ eq.	1.79×10^0	2.73×10^{-1}
Smog	kg O ₃ eq.	1.30×10^{-1}	2.27×10^{-2}
Acidification	kg SO ₂ eq.	1.57×10^{-2}	1.11×10^{-3}
Eutrophication	kg N eq.	3.93×10^{-3}	5.56×10^{-4}
Carcinogenics	CTUh	5.73×10^{-8}	1.54×10^{-7}
Non-carcinogenics	CTUh	1.39×10^{-7}	6.14×10^{-8}
Respiratory effects	kg PM2.5 eq.	1.41×10^{-3}	5.19×10^{-4}
Ecotoxicity	CTUe	4.77×10^0	6.02×10^0
Fossil fuel depletion	MJ surplus	2.12×10^1	2.14×10^{-1}

Table 8 LCA impacts for hydroelectric-powered processes due to varying DAC construction by −50% (low construction) and +50% (high construction), per kg of CO₂ captured and converted to methanol

Impact category	Unit	MW – low construction	MW – high construction	WH – low construction	WH – high construction
Ozone depletion	kg CFC-11 eq.	-3.61×10^{-9}	-2.21×10^{-9}	2.19×10^{-8}	2.31×10^{-8}
Global warming	kg CO ₂ eq.	-2.54×10^0	-2.52×10^0	-2.25×10^0	-2.23×10^0
Smog	kg O ₃ eq.	-1.94×10^{-2}	-1.80×10^{-2}	-1.17×10^{-2}	-1.05×10^{-2}
Acidification	kg SO ₂ eq.	6.21×10^{-4}	7.75×10^{-4}	1.43×10^{-3}	1.55×10^{-3}
Eutrophication	kg N eq.	-5.68×10^{-4}	-4.75×10^{-4}	-3.10×10^{-4}	-2.42×10^{-4}
Carcinogenics	CTUh	9.83×10^{-7}	9.96×10^{-7}	9.82×10^{-7}	9.92×10^{-7}
Non-carcinogenics	CTUh	1.29×10^{-6}	1.30×10^{-6}	1.31×10^{-6}	1.32×10^{-6}
Respiratory effects	kg PM2.5 eq.	-9.65×10^{-4}	-9.35×10^{-4}	-8.83×10^{-4}	-8.60×10^{-4}
Ecotoxicity	CTUe	1.17×10^2	1.20×10^2	1.16×10^2	1.17×10^2
Fossil fuel depletion	MJ surplus	-5.37×10^{-1}	-5.21×10^{-1}	-1.01×10^{-1}	-8.91×10^{-2}

Sensitivity analysis

Due to unknowns in the construction inventories, sensitivity analysis was performed per DOE guidance^{44,47} by varying the DAC construction inventories by $\pm 50\%$. Table 8 shows the impacts for each DAC regeneration heat source with hydroelectric power, since hydroelectric demonstrates the least environmental burden of all cases examined, and therefore is most sensitive to changes in construction. Results demonstrate that GWP varied by less than 1%, which substantiates previous studies that construction of DAC generally has a low life cycle impact when compared to the lifetime of operation energy use.⁸⁵ Ozone depletion varied the most by up to 27% in the MW case due to the high ozone burden of the MW inventory. Other impact categories were generally between 1–12% of the baseline results in Table 5.

Interpretation

Assessment of GWP impact

The results of this study indicate a significant potential for carbon removal, aligning with the objectives of global carbon neutrality goals. Notably, energy source and energy efficiency with capture technology were found to be the most critical in determining the life cycle carbon emissions and energy efficiency of these systems. Analysis reveals that DAC-to-methanol systems utilizing renewable energy sources exhibit markedly lower life cycle GHG emissions compared to those powered by conventional fossil fuels and energy mixes, as observed for DAC systems by previous studies.^{17,18,21,22} This stresses the importance of integrating DAC with sustainable energy systems to maximize environmental benefits. However, this study advances the conversation by providing a nuanced understanding of how specific low-carbon energy sources can optimize the performance and sustainability of DAC systems. This contributes valuable insights into the ongoing efforts to develop scalable and economically viable solutions for atmospheric carbon reduction. Future DAC research should focus on optimizing these systems for broader implementation, including exploring innovative regeneration energy solutions and sorbent technologies.

The main findings indicate that the most detrimental energy configurations for climate change are grid or fossil-fuel based



sources, as did McQueen *et al.*⁸⁶ who found that the integration of renewable energy sources with a DAC-based system provides profound environmental implications over natural gas. The GWP impacts of each energy scenario, shown in Fig. 6, highlight the substantial benefits of integrating renewable energy sources with synthetic fuel production and DAC technologies. Scenarios utilizing solar PV energy in AL and AZ demonstrate an improvement in GWP contributions compared to traditional sources but are impacted greatly due to the heavy construction and EOL environmental burden of PV infrastructure. The integration of hydroelectric, wind, and geothermal energy sources with green methanol and DAC technologies significantly lowers the GWP when compared to the US grid, further reinforcing that renewable energy sources are pivotal to the environmental sustainability. This research agrees with the study by Khojasteh-Salkuyeh *et al.*,¹² which found that the CO₂ hydrogenation process used for methanol production can be environmentally friendly but should be assessed on a regional level, and only where low-carbon energy sources are available.

The variability in environmental impacts across different DAC heat configurations highlights the importance of technology selection. The results indicate that scenarios employing WH recovery exhibit low GWP contributions, highlighting the efficiency and environmental benefits of leveraging WH in DAC processes in agreement with the study conducted by Deutz

and Bardow.⁴⁰ This suggests that optimizing DAC technology selection, based on environmental impact assessments, is crucial for minimizing the carbon footprint of methanol production systems. Research to further maximize the efficiency of MW technology for DAC regeneration demonstrates the life cycle improvements this technology can have on overall impacts, and continued research into this area is warranted.

The study's outcomes have profound implications for the implementation of DAC and carbon utilization technologies. By demonstrating the environmental advantages of renewable energy integration and the critical role of technology selection, the findings of this study provide a solid foundation for policies that encourage the adoption of sustainable green energy systems. The variability in GWP contributions across different scenarios brings to light the necessity for region-specific strategies that leverage local renewable energy resources, optimizing the sustainability of DAC-integrated methanol production. This study points to the need for further investigation focusing on optimizing DAC processes and technologies, particularly in the context of exploring new methods to improve energy efficiency and implementing new MW regeneration technology or WH recovery with renewable energy sources.

Assessment of other impacts

Traditional methanol production *via* SMR shows substantially higher impacts compared to any of the renewable energy

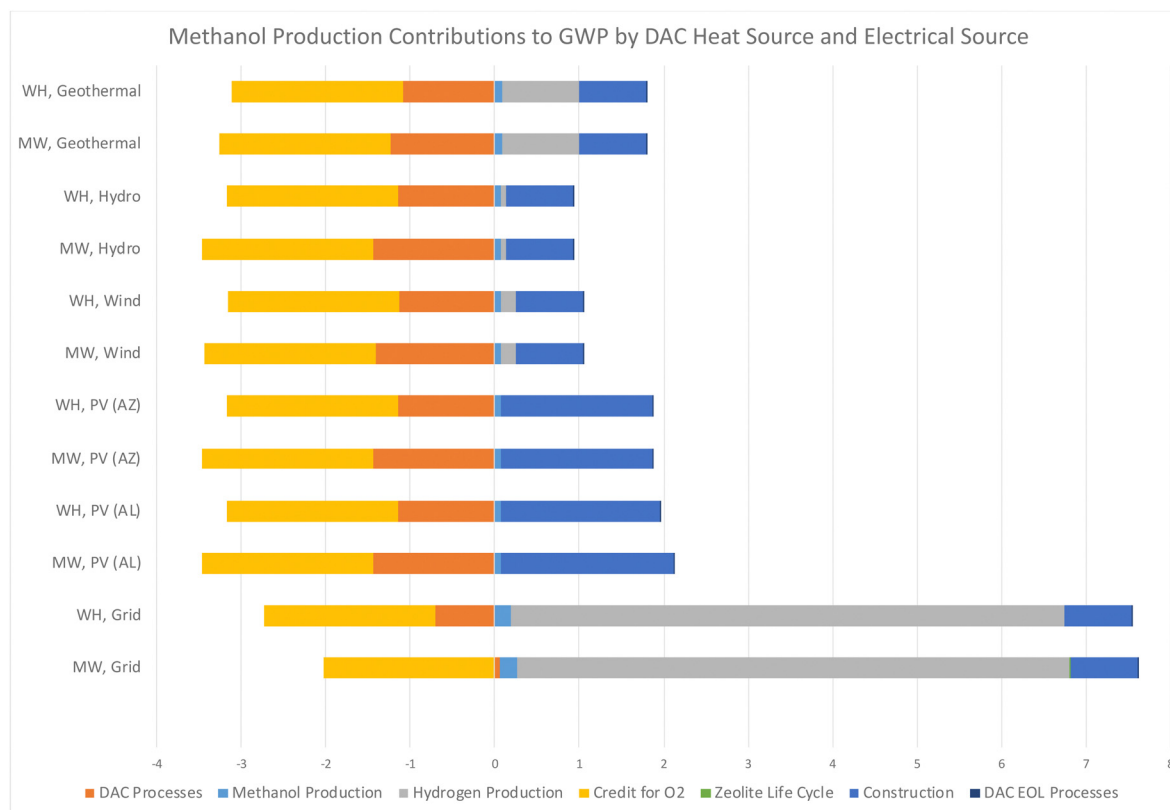


Fig. 6 Contributions to GWP for each configuration of a DAC regeneration heat source and electrical energy supply, per kg of synthetic methanol produced, in units of kg CO₂ eq.



The exploration of novel MW-based regeneration techniques, as compared to industrial WH, presents an innovative avenue for reducing energy consumption and improving system efficiency. Moreover, the integration of these systems with green hydrogen production, derived from water electrolysis powered by renewable energy, further enhances the environmental sustainability of the DAC-to-fuel process. This new

The scalability and infrastructure requirements for deploying DAC-to-fuel systems such as synthetic methanol are significant yet surmountable challenges that hinge on strategic integration

regeneration heat method should be targeted for future research to improve on the scalability to meet the requirements of modern, full scale DAC systems when industrial WH may not be an option.

Policy and market dynamics

In 2021, the global methanol market was valued at over \$37.4 billion and is projected to reach nearly \$61.7 billion by 2030, with a continuous annual increase in production worldwide.⁹⁰ In 2022, global methanol production was estimated to have exceeded 111 million metric tons, growing by nearly four percent from the previous year. The United States plays a significant role in the methanol market, both as a top exporter and a major importer. While methanol plays an important role in the US and world economies, it is important to shift toward more sustainable production and less dependence on fossil fuels to mitigate climate change.

Incorporating DAC systems and synthetic fuel production into the US market and with the current policy landscape is a crucial step toward mitigating climate change through carbon neutrality. This shift is reinforced by substantial federal support such funding allocations in legislation such as the Infrastructure Investment and Jobs Act,⁹¹ which has earmarked \$3.5 billion for carbon management technologies, and the Inflation Reduction Act of 2022,⁹² which has devoted an estimated \$370 billion total to energy security and climate change initiatives. Tax incentives such as the expanded 45Q tax credit⁹² further demonstrate the financial viability of DAC projects. These policy frameworks are complemented by dynamic market forces, where both start-ups and established corporations are venturing into DAC and synthetic fuels, spurred by potential carbon offset markets and sustainability objectives. Collaborations across private, governmental, and academic spheres are crucial for surmounting technical hurdles and scaling solutions. Yet, the commercial deployment of DAC technologies and synthetic fuel production is fraught with challenges, chiefly cost and energy efficiency. Innovations aimed at reducing these barriers are vital for competitive parity with traditional fuels and carbon capture methods, marking a pivotal era for DAC systems and synthetic fuel production in aligning economic incentives with climate goals.

Comparison to other studies

While the literature on methanol production *via* point source liquid amine based CCU is extensive, DAC integrated with methanol synthesis is less commonly studied. This discrepancy limits the possibility of direct comparisons between the two methods. However, understanding the differences in GHG impacts between traditional CC and innovative DAC methods is crucial for evaluating the potential and efficiency of newer technologies like DAC in achieving lower emissions. Therefore, the results for GWP from this study were compared to the assessment by Cordero-Lanzac *et al.*,²⁹ who compared the impacts of green methanol production from amine-based point source carbon capture with two different energy configurations: renewable and non-renewable sources in the EU. Their results,

converted to this study's functional unit, found that renewables demonstrated a GWP impact of about -1.45 kg CO₂ eq. while non-renewables impacted by about 1.36 kg CO₂ eq. These results demonstrate, similarly to this study, the potential for CO₂ reduction *via* using renewably powered processes.

This study aligns well with the findings of Galusnyak *et al.*,³¹ who performed an LCA of methanol production and conversion into various chemical intermediates and products. Both studies highlight the significant influence of energy sources on environmental impacts. Galusnyak *et al.* assessed three energy scenarios: EU grid mix, wind power, and hydro power. They found that using renewable energy sources substantially reduces environmental impacts. Similarly, our study demonstrates that hydroelectric power has the lowest impacts across various categories, while the US grid mix shows the highest impacts. For GWP, Galusnyak *et al.* reported substantial reductions when using wind and hydro power compared to the grid mix, aligning with our findings where hydro and wind exhibit the lowest GWP. Both studies emphasize the importance of renewable energy in achieving better environmental performance for methanol production and highlight the trade-offs involved in different energy sources.

Our findings are also compared to Kibria Nabil *et al.*,⁴⁵ which performed a comparative LCA of electrochemical upgrading of CO₂ to fuels and feedstocks. The authors highlighted that the carbon intensity of electrochemical routes is highly sensitive to the electricity emission intensity, achieving climate benefits only when coupled with low-emission electricity (<0.25 kg CO₂ eq. per kW h). Similarly, our study found that the US grid mix results in the highest impacts, whereas renewables show the lowest. For other impact categories, both studies report that electrochemical processes can have significant energy requirements, particularly in conversion and separation phases, which can offset climate benefits unless powered by renewable energy sources. Our findings of high impacts associated with PV systems in Alabama, for example, mirror the authors' emphasis on the need for renewable optimization to ensure overall environmental benefits. Overall, both studies highlight the potential of renewable energy-based systems to substantially reduce GWP and other environmental impacts, while also emphasizing the challenges and trade-offs involved in optimizing these technologies.

Similarly, this study's results are compared to Maimaiti *et al.*,³⁶ who examined renewable energy-based methanol production in China. Both studies highlight the critical influence of energy sources on environmental impacts, with hydroelectric and wind energy showing the lowest impacts. Maimaiti *et al.* reported a GWP of 0.105 kg CO₂ eq. per kg methanol for a photovoltaic-based system with battery storage, similar to our findings where hydroelectric and wind power exhibit the lowest impacts. Both studies also show significant reductions in fossil fuel depletion and GWP for renewable systems compared to conventional methods. However, both studies highlight higher impacts for photovoltaic systems in human toxicity and ozone depletion due to polycrystalline silicon production. This study corroborates Maimaiti *et al.*'s findings, demonstrating the



technology integration, emphasizing the promise of DAC with green methanol production as a sustainable path towards achieving carbon neutrality and fostering a greener future.

Nomenclature

Al	Aluminum
AL	Alabama, USA
AZ	Arizona, USA
C	Celsius
CCU	Carbon capture and utilization
CFC-11	Trichlorofluoromethane
CO ₂	Carbon dioxide
CTUe	Comparative toxic unit for ecosystems
CTUh	Comparative toxic unit for health
Cu	Copper
DAC	Direct air capture
DOE	United States Department of Energy
EOL	End of life
eq	Equivalent
EPA	United States Environmental Protection Agency
EU	European Union
FU	Functional unit (of LCA)
GHG	GHG
GW h	Gigawatt-hour
GWP	Global warming potential
h	Hour
IPCC	Intergovernmental Panel for Climate Change
kg	Kilogram
kt	Metric kiloton
kW	Kilowatt
kW h	Kilowatt-hour
LCA	Life cycle assessment
LCI	Life cycle inventory analysis
LCIA	Life cycle impact assessment
m	Meter
min	Minute
mL	Milliliter
MJ	Megajoule
MSD	Microwave swing desorption
MW	Microwave
N	Nitrogen
NET	Negative emissions technology
NO _x	Nitrogen oxides
O ₂	Oxygen
O ₃	Ozone
PEM	Polymer electrolyte membrane
PM2.5	Particulate matter <2.5 microns
PV	Solar photovoltaic
SO ₂	Sulfur dioxide
SMR	Steam methane reforming
Ton	Metric ton
TRACI	Tool for reduction and assessment of chemicals and other environmental impacts
VOC	Volatile organic compound

This study's findings underscore the critical importance of strategic technology selection and the optimization of energy sources in minimizing environmental impacts and promoting sustainable energy solutions. By showcasing the environmental benefits of integrating renewable energy with DAC and methanol production technologies, this research contributes valuable insights to the ongoing efforts in scaling up these technologies for global climate mitigation.

As the world seeks viable solutions to combat climate change, the integration of DAC with green methanol production emerges as a promising pathway towards carbon neutrality, offering a significant step forward in the quest for sustainable and environmentally friendly energy solutions. This study lays a foundation for future research and policy development, aiming to accelerate the deployment of DAC and renewable energy technologies and thereby contributing to the overarching goal of mitigating global warming and achieving a sustainable future. By presenting a comprehensive analysis of renewable energy scenarios for DAC-to-methanol synthesis, the study contributes a significant chapter to the body of knowledge necessary for the advancement of these technologies towards global climate mitigation efforts. The findings advocate for a nuanced approach to

W	Watt
WECC	Western Electricity Coordinating Council
yr	Year
Zn	Zinc

Data availability

This paper was written based on publicly available information, and all values used in calculations are explained or cited. An extensive life cycle inventory and detailed results are available in the ESI.†

Conflicts of interest

There are no conflicts of interest to declare.

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