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Green ethylene production in the UK by 2035: a techno-economic assessment†

Andreas H. Nyhus,‡ Maria Yliruka, *‡ Nilay Shah  and Benoît Chachuat *

Olefins production in the UK is the most emission-intensive sector of the chemical industry. Bringing thermocatalytic and electrocatalytic processes together, this paper compares nine process routes for green ethylene production from air-captured CO₂ and off-shore wind electricity in order to displace fossil-based ethylene, with a particular focus on technology readiness for near-future deployment. The methanol-mediated thermocatalytic route has the lowest projected levelised cost at £2900 per ton of ethylene by 2035, closely followed by direct and tandem CO₂ electroreduction routes in the range £2900–3200. The price of green ethylene at three times or more its current market price is confirmed through a sensitivity analysis varying the levelised cost of electricity, stack cost, and market price of propylene or oxygen simultaneously. While these green ethylene production processes would be carbon negative from a cradle-to-gate viewpoint, displacing a conventional ethane cracker with annual production capacity of 800 kt could consume as much as 46–66 TW h of renewable electricity, which is a major barrier to deployment.

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Broader context

Ethylene is the most widely produced organic compound globally. For decades, steam cracking has been the dominant method to produce ethylene from fossil fuels but its production is one of the largest CO₂-emitting processes in the chemical industry, calling for the development of more sustainable production routes. Of the alternatives to defossilise ethylene, is the use of CO₂ captured from ambient air as feedstock and renewable electricity from offshore wind in order for the produced ethylene to be fully compliant with the green chemical standard. Candidate technologies for producing such green ethylene at scale include a range of conventional and unconventional, thermocatalytic and electrocatalytic processes, which have to date been assessed separately and often using process boundaries that fail to comply with the green production standard. The techno-economic assessment here fills in this gap and seeks to predict a lower bound on future market price of green ethylene by 2035, with a particular focus on the UK context.

1 Introduction

Light olefins, comprising ethylene and propylene, are fundamental building blocks in the chemical industry for the production of plastics, synthetic fibers and synthetic rubbers.^{1,2} In Europe light olefins are mostly produced through naphtha steam-cracking, while in the United Kingdom (UK) and the United States (US) alkane steam crackers are more common.^{3–5} A typical ethane cracker produces about 50% ethylene, 35% ethane, 5% C₃₊, and additional light hydrocarbons.² In the UK alone, the annual olefins production amounts to 2.75 Mt and emits 3.1 Mt of CO₂, making it the most emission-intensive

process across the UK's chemical sector (share of 17%).^{6,7} It contributed about 0.5% of the UK's total greenhouse gas (GHG) emissions in 2019.^{8,9} Defossilising olefins production is particularly challenging given that fossil fuels are not only used to meet the utility demands, but also as the carbon feedstock itself.¹⁰

Biomass, polymers and CO₂ are all candidate feedstocks for defossilising light olefins and olefin-derived products.^{11–17} Since sustainably-sourced lignocellulosic biomass is in limited supply, its use may rather be prioritised for negative emission technologies such as bioenergy for power plants coupled with carbon capture and long-term geological storage.¹⁸ Challenges with polymer recycling, either mechanically or chemically, arise due to feedstock heterogeneity and availability.¹⁹ The number of times a polymer can undergo mechanical recycling is furthermore limited due to the degradation of its mechanical properties,²⁰ while additives and impurities can cause catalyst deactivation in chemical recycling processes.^{21,22} In contrast to

The Sargent Centre for Process Systems Engineering, Department of Chemical Engineering, Imperial College London, Exhibition Road, London, SW7 2AZ, UK.
E-mail: m.yiliruka@imperial.ac.uk, b.chachuat@imperial.ac.uk

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‡ These authors contributed equally to this work.



biomass and plastic waste, CO₂ captured from industrial processes or extracted from the atmosphere can be purified to meet process specification for its utilisation. The UK has large CO₂ storage capacities and aims to become a global leader in carbon capture, utilisation, and storage.²³ The industrial clusters around Merseyside and on the East coast are expected to capture, respectively, 10 and 27 Mt_{CO₂} annually from industrial point sources from 2030.^{24–26} Large quantities of CO₂ could therefore become available for CO₂ utilisation projects, although no such development is yet planned in the UK.²⁷ While CO₂ from industrial processes and power plants remains ultimately linked to fossil fuel consumption, direct air capture (DAC) already has a negative carbon footprint for the current UK electricity mix (0.18 kg_{CO₂} kW h⁻¹ in 2022).^{28,29} Following the terminology applied to hydrogen production processes,³⁰ only a production route based on atmospheric carbon and renewable energy may indeed qualify as green.

Two pathways for the utilisation of CO₂ using renewable energy include: (i) thermocatalytic processes reliant on green hydrogen, and (ii) electroreduction of CO₂ (CO₂RR) powered by renewable electricity. The former can benefit from the economy of scale for large production volumes, minimize utility demands *via* heat integration, and produce valuable hydrocarbon co-products. Although at a lower technology readiness level (TRL) than thermocatalytic processes, CO₂RR avoids losses incurred in converting electricity to hydrogen and will benefit from the large technological learning rates projected for hydrogen electrolyzers in the near future (*e.g.*, from \$880 kW⁻¹ today down to \$400 kW⁻¹ by 2030 for 10 MW PEM electrolyzers).³¹ CO₂RR may be used to directly produce ethylene or alternatively produce an intermediate that is subsequently converted to ethylene. Nevertheless, low-margins force the chemical industry to be risk-averse, causing a preference for thermocatalytic technologies which have greater maturity.³² Additionally, the delay induced by waiting for a low-TRL technology to become mature comes at the risk of missing emission reduction targets.³³ There is a need, therefore, for comparing the available routes from CO₂ and renewable energy to ethylene in a systematic manner, through taking current performance into account to decide whether to prioritise investment into scale-up or further research-and-development. Investment into the defossilisation of ethylene would have a significant impact across the chemical sector, as its embedded emissions are currently a major contributor to the cradle-to-gate emissions of numerous downstream products, including low- and high-density polyethylene (LDPE, HDPE), polyethylene terephthalate (PET), and polyvinyl chloride (PVC).^{34,35}

1.1 Techno-economic review

Multiple techno-economic assessments of CO₂ electroreduction processes have set out to determine which products could become competitive with conventional production routes, most commonly, formic acid, ethanol, carbon monoxide and ethylene.^{36–39} Orella *et al.*³⁷ further included methane, Verma *et al.*³⁸ methanol, and Jouny *et al.*³⁹ methanol and *n*-propanol. Having the highest product value per electron, formic acid and

carbon monoxide were the only products with positive end-of-life net present values under current market prices. Ethylene produced *via* CO₂RR could be competitive with naphtha cracking, although not with ethane cracking.³⁶ None of these studies focused on green ethylene production but instead assumed processes powered with grid electricity in a price range of \$30–60 MW h⁻¹ and fed with CO₂ captured from large point sources at a price of \$16.5–70 t_{CO₂}⁻¹ to minimise production cost. Disregarding the alkaline conditions commonly used in CO₂RR experiments, idealised assumptions were made for CO₂ electrolyzers with proton-exchange membrane (PEM) in terms of their faradaic efficiency, current density, and single-pass conversion. Additionally, liquid and gas separation costs were taken proportional to the product flowrates rather than based on the actual product composition.

Going beyond the CO₂ electroreduction processes, existing comparisons between thermocatalytic and electrocatalytic processes for producing methanol,^{40,41} methane⁴² and syngas⁴³ from CO₂ concluded that the former would be more profitable. It was also found that the production cost of these chemicals is typically driven by operating costs, accounting for more than 60% of the total cost. The losses incurred in the conversion of renewable electricity to H₂ in any of the thermocatalytic routes could be afforded because water electrolyzers presently enjoy higher energy efficiencies (around 70%) than CO₂ electrolyzers (dependent on target product but typically below 50%).^{40,42,43} A limitation in all of these studies is that the levelised cost of renewable electricity was set equal to the price of electricity, thereby neglecting the cost of balancing both daily and seasonal intermittency. Kim *et al.*⁴¹ even assumed a plant operation using free, surplus renewable electricity. Again, idealised assumptions on the performance of CO₂ electrolyzers were made,^{40,41} and only Welch *et al.*⁴² and Moreno-Gonzalez *et al.*⁴³ considered a CO₂ feedstock from DAC, which is necessary for the produced chemicals to qualify as green.

Regarding the complete route from CO₂ to ethylene using renewable electricity, the thermocatalytic conversion of CO₂ to methanol then to olefins (MS+MTO) was compared to CO₂ methanation followed by oxidative coupling of methane (CTM+OCM)⁴⁴ and to direct Fischer–Tropsch synthesis from CO₂ (D-FT).⁴⁵ Based on the energy content of the products, ethylene from D-FT could cost eight times as much as from MS+MTO due to the large utility requirement for the compression and pre-heating of its recycle stream. Ethylene from CTM+OCM could also be twice as costly as from MS+MTO,⁴⁴ although the corresponding assessment did not account for differences in the product stream composition. In separate comparisons,^{46,47} both the MS and MTO stages were found to be cheaper than CTM and OCM, respectively. The cost advantage of MS over CTM was driven by the lower H₂:CO₂ ratio while MTO out-competed OCM by generating a larger revenue from selling propylene compared to ethylene in the US market. Focusing only on electrocatalytic processes, Ramdin *et al.*⁴⁸ compared the direct CO₂RR to ethylene (D-eC₂H₄) to a two-stage electrolysis process converting CO₂ to CO either at high- or low-temperature, followed by CO electroreduction (CORR) to ethylene. The high-



temperature two-step route was predicted to be cheaper, while the total cost of both low-temperature routes were comparable. Modelling the separation processes in more detail than other studies, Ramdin *et al.*⁴⁸ also confirmed the dominant costs to be the capital and operating costs of the electrolysers, contributing over 75% of the total system cost. Bringing together both types of processes, Ioannou *et al.*⁴⁹ compared the electrocatalytic D-eC₂H₄ route to the thermocatalytic MS+MTO route. The latter was found to be the cheaper alternative since it had lower capital costs and could benefit from the higher energy efficiency of H₂ electrolysers (65%) compared to CO₂ electrolysers (34%). For scenarios in which electricity was generated by nuclear reactors or onshore wind turbines, Ioannou *et al.*⁴⁹ predicted negative global warming potentials for both routes, yet production costs two to three times larger than naphtha cracking.

1.2 Contributions and outline

The techno-economic review above identifies the methanol-mediated thermocatalytic route as a promising candidate for low-cost green ethylene production from CO₂ and renewable energy. While combined electrocatalytic and thermocatalytic processes *via* syngas and methanol have been reviewed,⁵⁰ none of these combinations have been systematically compared through techno-economic analysis. Disparate system boundaries and costing methodologies in existing, often pairwise assessments make it difficult to draw meaningful conclusions about their overall cost-competitiveness.

To supply either CO₂ utilisation pathway with green electricity and hydrogen, the UK's renewable power generation capacity would need to be expanded.⁵¹ A whole systems perspective is much needed to account for the limitations on renewable generation capacity due to land availability, cost and public acceptance.⁵¹ However, current scenarios for the UK's future energy system fail to reflect these additional demands,^{52–54} due in part to the difficulty of integrating chemical production processes into whole-system energy modelling. Deciding early on a decarbonisation strategy across different sectors is however paramount to reducing the overall system costs associated with the transition to net-zero by 2050.⁵⁵

This paper presents a comprehensive techno-economic analysis of nine process routes for green ethylene production from air-captured CO₂ and renewable electricity from a dedicated offshore wind farm, by bringing thermocatalytic and electrocatalytic processes together. Using consistent system boundaries, process inventories are derived from best-in-class, heat-integrated thermocatalytic processes and both low- and high-temperature CO₂ or CO electrolysers. The candidate routes are ranked according to the levelised cost of ethylene and a sensitivity analysis is conducted to confirm that the conclusions are robust towards changes in the levelised cost of electricity, stack cost, and market price of co-products. While the focus of this work lies in the comparison of defossilised ethylene production pathways on a process level, the process inventories computed herein could be used as part of any energy system model too.

The rest of the paper is structured as follows: Section 2 starts with reviewing the technology readiness and process conditions

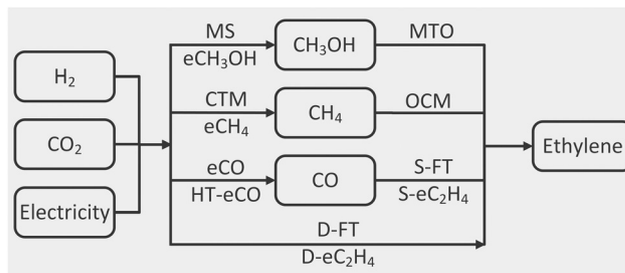


Fig. 1 Overview of the processes and intermediates in the conversion of CO₂ to ethylene. Abbreviations: MS: methanol synthesis; eCH₃OH: CO₂ electroreduction to methanol; MTO: methanol to olefin; CTM: CO₂ methanation; eCH₄: CO₂ electroreduction to methane; OCM: oxidative coupling of methane; eCO: CO₂ electroreduction to CO; HT-eCO: high-temperature CO₂ electroreduction to CO; S-FT: syngas Fischer–Tropsch; S-eC₂H₄: CO electroreduction to ethylene; D-FT: direct Fischer–Tropsch; D-eC₂H₄: CO₂ electroreduction to ethylene. Refer to Section A of the Appendix for details.

for selected production routes to green ethylene, then details the process modelling and levelised cost calculation methodology. Section 3 discusses the predicted process inventories and levelised costs of ethylene production for each route, before analysing the sensitivity of the top two routes, and examining systemic consequences beyond the process boundaries. Finally, Section 4 concludes the paper.

2 Methodology

2.1 Technology selection and modelling

A common system boundary is set around each process so that only renewable electricity, green hydrogen, CO₂, air and water can enter, while only ethylene, hydrocarbon co-products, oxygen, residual CO₂ and wastewater can leave. All hydrocarbon mixtures are separated in terms of their ethylene, ethane, propylene, propane, methane or C₄₊ constituents. Residual emissions from purge streams and carbon-capture processes and wastewater from the separation and purification are also accounted for as outputs.

The processes of interest (Fig. 1) convert CO₂ into ethylene either directly or *via* a single one-carbon intermediate (CO, CH₄ and CH₃OH). The conversion of CO₂ to an intermediate is referred to as stage-1 and the subsequent conversion of the intermediate to ethylene as stage-2. For those processes needing syngas in stage-2, *in situ* produced CO is combined with green H₂ in the relevant stoichiometry ratio. The ethylene production rate from stage-2 governs the production rate of the intermediate from stage-1, and both stages are integrated. Any unconverted CO₂ or H₂ from stage-2 is recycled to stage-1. Similarly, the heat and electricity utility demands of one stage may be met by excess heat or electricity from the other stage. A more systematic process integration using optimisation was beyond the scope of this work.

A key outcome of the modelling conducted herein is linear input-output models for each (single- or two-stage) process derived from the detailed mass and energy process inventories



and normalised to the production of 1 tonne of ethylene in the manner of a functional unit in life-cycle assessment (LCA). Such linear models are particularly suitable for integration into energy system models as part of future research.

2.1.1 Thermocatalytic processes. Three criteria were applied for the selection of thermocatalytic process models: first, a reactor model should be based on reaction kinetics to reflect realistic performance levels, with preference given to commercial processes where available. Second, a process should be heat-integrated in order to minimise utility requirements and reflect the true potential of thermocatalytic processes. Third, the flowsheet, mass and energy balances, and cost analysis should be explicitly modelled to enable rescaling of all the relevant units under a common system boundary.

The reference process are adapted to the new system boundary for green ethylene production by excluding any process unit generating an intermediate from fossil resources and including an air separation unit where a pure oxygen feed stream is required.⁵⁶ Where applicable, the cryogenic distillation train by Liu *et al.*⁵⁷ is added for the separation of hydrocarbon mixtures.

The mass and energy balances are subsequently updated, including the recycling of concentrated CO₂ and H₂ streams and any purge and flue gas streams. The utility demands are met by first integrating any excess heat and electricity. Where a net heat surplus is available after integration, a condensing steam turbine is designed to produce electricity.⁵⁸ And where a net import of hot utility is required, the system is extended to include a hydrogen boiler. Finally, capital expenses from the reference processes are scaled according to the six-tenth rule⁵⁸ and converted to 2019 prices using the Chemical Engineering Plant Cost Index (CEPCI).⁵⁹

Refer to Section A of the appendix and Section S1 of the ESI,[†] for further details about the reference processes, including any adjustments made.

2.1.2 Low-temperature electrocatalytic processes. Although low-temperature CO₂ and CO electrolyzers have improved tremendously over the last decade, their technological maturity remains low (TRL 4). The screening of electrocatalytic processes therefore focuses on state-of-the-art lab-scale performance with preference given to: common cathode material, membrane type and electrolyte; flow cell experimental set-ups over H-cell configurations;⁶⁰ and experimental studies showing sustained performance for longer than 1 hour. Where several such studies can be found, the final choice is based on the highest current density (to reduce the capital cost) and lowest cell voltage (to reduce the electricity demand).³⁹

The electrochemical model describes the relationship between the electrical power and CO₂ (or CO) feed rate with the flow rate of product. The electric current is set by the flow rate and the faradaic efficiency (η_F) of the product, while the required electrode area (A_{cell}) follows from the current density (j). The cell voltage (E_{cell}) is calculated based on the half-cell potentials of the CO₂RR (E_{red}) and oxygen evolution reaction (OER), which depart from the standard cell potentials due to both activation potentials and ohmic losses. While the half-cell

Table 1 Parameters used in the electrochemical model of each reference process. Refer to Appendix A for details about the reference processes and to Section S2.1.1 of the ESI for cell voltage calculations

Process	Ref.	Parameters				
		η_F (—)	X (—)	j (A m ⁻²)	E_{red} (V)	E_{cell} (V)
eCO	69	1.00	0.40	3500	1.63	2.83
eCH ₄	70	0.73	0.40	5000	2.31	3.51
S-eC ₂ H ₄	71	0.69	0.40	1500	1.61	2.81
D-eC ₂ H ₄	72	0.66	0.40	1100	1.43	2.63

potential at the anode could be reduced by oxidising organic molecules instead of water,^{61,62} this option would not be compliant with the set system boundary around renewable electricity, H₂, CO₂, air and water (*cf.* Section 2.1). The flow rates of the co-products and reactants are estimated from the product flow rate, the electrochemical reaction's stoichiometry (*cf.* Section A of the Appendix) and the single-pass CO₂ or CO conversion (X). The parameter values for the electrocatalytic processes of interest are summarised in Table 1. The faradaic efficiencies, current densities, and cell voltages were all measured experimentally. The set single-pass CO₂ or CO conversions of 40% were not measured, but deemed achievable by 2035 based on 10–40% conversion values reported in recent alkaline zero-gap electrolyzers,^{63–66} largely limited by carbon losses due to carbonate formation.^{67,68}

For the scale up of CO₂ and CO electrolyzers, a zero-gap cell design is preferred to a flow cell design as it reduces the overall cell resistance and increases performance, particularly at high current densities.^{66,73} The cell design and the balance of plant (BOP) layout of anion-exchange membrane (AEM) electrolyzers are analogous to those of PEM electrolyzers.⁷⁴ A similar expression therefore is used for their capital cost, building on a specific stack cost estimate around £280 kW⁻¹ by 2035.⁷⁵ For 1 MW PEM electrolyzers, the stack cost contributes about 35% of the total system cost;⁷⁶ the BOP makes the other 65% in terms of power supply, deionised water circulation system, gas processing and cooling equipment, all expected to be manufactured at scale for electrolyser systems by 2035. Whereas the cost of the BOP scales with the power consumption of the electrolyser, the stack cost correlates with the electrode area.

To separate ethylene from the mixed gas products of the electrolyzers, vacuum pressure swing adsorption (VPSA) processes are used,^{48,77} and any unconverted CO₂ is recovered and recycled at a 97% capture rate by mixed-amine CO₂ absorption.⁷⁸ The separation of the liquid co-products (acetic acid, alcohols) is also considered when their concentration in the electrolyte exceeds 20 wt%. The utility demands of the separation train are met by hydrogen boilers and renewable electricity imports because they cannot be heat-integrated with the low-temperature electrolyzers. Both the capital and utility costs of the separation processes are scaled up according to the total product flow rate. The total invested capital (TIC) comprises the fixed capital investments for the CO₂ electrolyser and its associated separation train, as well as additional working capital, start-up and contingency costs.⁷⁹



Refer to Section A of the Appendix for additional details about the reference CO₂ electrolyzers and to Section S2 of the ESI,[†] for further details about the electrochemical model and the costing methodology for the low-temperature CO₂ electrolyzers and their separation units.

2.1.3 High-temperature electrocatalytic processes. High-temperature, solid-oxide electrolysis cells (SOEC), including dry electrolysis for converting CO₂ into CO, have undergone tremendous improvements over the past 15 years in terms of their electrochemical performance and long-term durability, both at the cell and stack levels.^{80,81} Rather than developing a bottom-up model based on lab-scale experimental data as is done for low-temperature electrolyzers (*cf.* Section 2.1.2), the focus is on Haldor Topsoe's carbon monoxide generator, eCOs™ for the SOEC modelling owing to its higher maturity (TRL 8).^{82,83} The model assumes an overall conversion of 100%, heat supplied by electric heaters, and gas purification processes (>99 vol% of CO) included in the total energy consumption (8 kW h per Nm³ of CO produced)—this is in contrast to the low-temperature electrolyser models where H₂ is consumed to supply heat to the amine absorption process for CO₂ recycling.

Based on a scenario in which the cumulative capacity continues to grow at the same rate as in the past, a specific cost of £600 kW⁻¹ is assumed for the SOEC system by 2035.⁸⁴ This represents a >70% reduction from the present cost of 1200–2000 kW⁻¹.^{84–88} The future capital cost of SOEC systems are perceived to be more uncertain than that of alkaline or PEM electrolyzers,⁸⁹ given the limited number of installations today⁸⁴ and the anticipated technological improvements.⁹⁰ The assumption that learning from high-temperature water electrolysis systems can be transferred to high-temperature CO₂ electrolysis is consistent with the low-temperature CO₂ electrolyzers above.

Refer to Section A of the Appendix for additional details about the reference SOEC electrolyzers and to Section S2 of the ESI,[†] for further details about the SOEC performance and costing.

2.2 Techno-economic analysis

The candidate process routes are compared in terms of the levelised cost of ethylene (LCOEt, eqn (1)), which accounts for the resource consumption, the revenues from co-products, and the annualised capital costs of the plant at scale. The cash-flow calculation assumes that the plant is erected during the first year, then operated at full capacity from the second year onward. The return on investment is neglected by setting the net present value of the project to zero.

$$\text{LCOEt} = \text{AF}_{\text{C}_2\text{H}_4} \frac{\text{OpEx} + \text{CapEx CRF}}{m_{\text{C}_2\text{H}_4}} \quad (1)$$

The capital expenditures (CapEx) are annualised using a uniform capital recovery factor (CRF). The operational expenditures (OpEx) comprise the variable utility costs and the fixed operating expenses for salaries, maintenance and overheads.

The economic allocation factors (AF_{*j*}, eqn (2)) apportion the total annualised cost of a process among all the co-products

Table 2 Nominal values of economic input parameters in the techno-economic analysis. The ranges provided for selected uncertain parameters in the sensitivity analysis assume triangular distributions. The value of Cost_{stack} is only applicable to AEM cells. The rows in bold indicate that the values of Cost_{el}, Cost_{H₂} and Cost_{CO₂} are varied simultaneously in the sensitivity analysis. The higher heating value (HHV) of hydrogen (141.8 MJ kg⁻¹)¹⁰⁶ is used

Parameter	Unit	Nominal	Range
LT	year	20	
<i>r</i>	—	0.06	
UF	—	0.9	
CEPCI ₂₀₁₉	—	607.3	
<i>m</i> _{C₂H₄}	kt year ⁻¹	800	
Cost _{tax}	£ t _{CO₂} ⁻¹	301.4	
Cost _{t_{ww}}	£ t _{ww} ⁻¹	0.52	
Cost_{el}	£ MW h ⁻¹	54.6	49.9–76.7
Cost_{H₂}	£ MW h ⁻¹	74.3	67.9–104.3
Cost_{CO₂}	£ t _{CO₂} ⁻¹	312.3	301.5–363.1
Cost _{stack}	£ kW ⁻¹	283.5	235–782
Ratio _{CH₄}	—	0.13	
Ratio _{C₂H₆}	—	0.15	
Ratio _{C₃H₆}	—	0.87	0.70–1.05
Ratio _{C₃H₈}	—	0.39	
Ratio _{C₄₊}	—	0.8	
Ratio _{O₂}	—	0.13	0.00–0.13

(*j* ∈ {CH₄, C₂H₄, C₂H₆, C₃H₆, C₃H₈, C₄₊, O₂}), according to their share on the overall revenue.

$$\text{AF}_j = \frac{m_j \text{Price}_j}{\sum_k m_k \text{Price}_k} \quad (2)$$

Green chemical products are expected to be sold at a price premium compared to their fossil-based counterparts. Herein, the prices of the co-products (Price_{*j*}) are assumed to increase alongside the price of ethylene (Price_{C₂H₄}), so their relative market prices (Ratio_{*j*}, eqn (3)) with respect to ethylene will remain constant throughout the plant's lifetime. The expression for AF_{C₂H₄} using these relative market prices is given in eqn (4).

$$\text{Ratio}_j = \frac{\text{Price}_j}{\text{Price}_{\text{C}_2\text{H}_4}} \quad (3)$$

$$\text{AF}_{\text{C}_2\text{H}_4} = \frac{m_{\text{C}_2\text{H}_4}}{m_{\text{C}_2\text{H}_4} + \sum_{j \neq \text{C}_2\text{H}_4} m_j \text{Ratio}_j} \quad (4)$$

Refer to Sections S4 and S5 of the ESI,[†] for further details on economic calculations.

2.3 Case study and sensitivity analysis

The ethane cracker in Wilton (North Yorkshire, UK), with an annual production capacity of 800 kt, is chosen as the reference herein.^{5,91} A plant lifetime of 20 years is assumed, with a utilisation factor (UF) of 90% and discount rate of 6%.^{92,93} To study a transition to green ethylene in the near future, cost assumptions for 2035 are used where available, with all costs expressed in £₂₀₁₉ for consistency. The economic input parameters are summarised in Table 2 and discussed below.



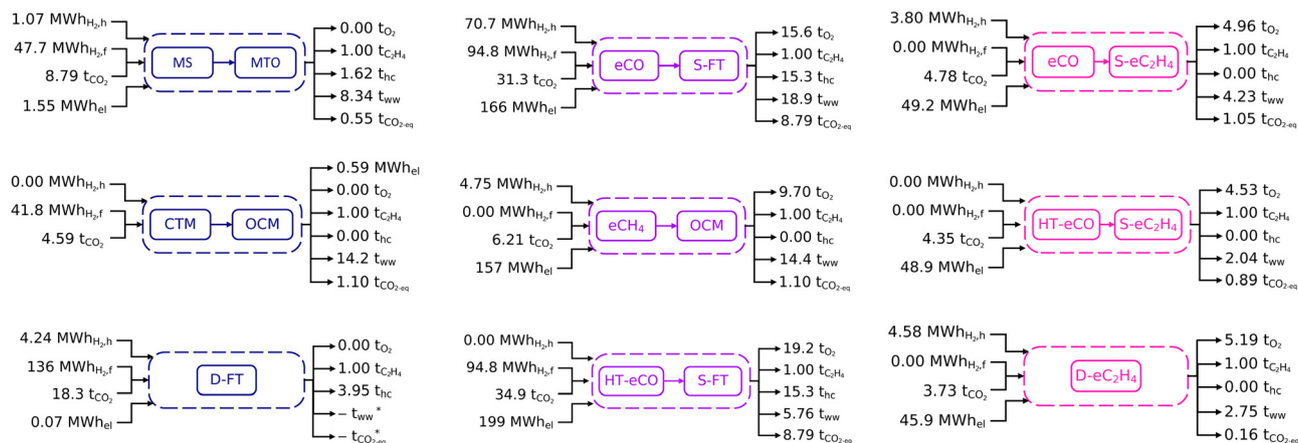


Fig. 2 Process inventories for the production of 1 t of ethylene using fully thermocatalytic routes (blue), fully electrocatalytic routes (pink) and hybrid routes (purple). The hydrogen inputs distinguish heat (H₂,h) and feedstock (H₂,f). The hydrocarbon by-products (hc) include methane, ethane, propylene, propane and C₄₊. The waste streams consist of wastewater (ww) by the separation processes and residual GHG emissions (CO₂-eq). *Not quantified.

While such large-scale production capacity is instrumental to contextualise the electricity consumption of green ethylene (*cf.* Section 3.4 below), some of the emerging technologies may not reach kilotonne annual production levels in the near future. It is worth reiterating that the process inventories and LCOEt for each route are calculated for smaller (relevant) process scales, as dictated by their reference processes (*cf.* Section A of the Appendix). The assessment methodology therefore relies both on technology scale-up and technology scale-out.

The cost of non-intermittent electricity (Cost_{el}), renewable hydrogen (Cost_{H₂}) and air-captured CO₂ (Cost_{CO₂}) are based on a common value for the levelised cost of electricity (LCOE) of offshore wind in the UK.⁷⁵ Balancing intermittency using both short- and long-term storage for the uninterrupted supply of renewable electricity is assumed to add 30% on top of the LCOE.⁹⁴ The price of domestically produced green H₂ is based on the LCOE using PEM electrolyzers with an efficiency of 1.3 MW h_{el} MW h_{H₂,HHV}⁻¹, and adding an extra 36% for capital and operating costs of the electrolyzers.⁷⁵ Given the proximity of underground salt caverns to the Wilton production site⁹⁵ and existing plans to build a hydrogen pipeline network in the North East of England by 2030,⁹⁶ additional charges for transport and storage are henceforth neglected.^{97,98} For the ethylene product to be considered green, only CO₂ from DAC is deemed a suitable feedstock. A cost of £312.3 t_{CO₂}⁻¹ is estimated on the assumption that solid sorbent-based DAC is deployed at scale by 2035, including automated manufacturing of the CO₂ collectors and based on the average relative humidity and temperatures in the UK,⁹⁹ which is six-times higher than the cost for CO₂ capture from cement.^{100,101} Air and water are assumed to be free of charge. The cost of residual process emissions (Cost_{tax}) is based on the 2035 UK carbon price of £301.4 t_{CO₂}⁻¹,¹⁰² while a generic cost of £0.52 per ton of treated wastewater (Cost_{ww}) is assumed.¹⁰³

A further assumption in the economic comparison is that all the hydrocarbon co-products can be sold separately, although they are often integrated with downstream processing or combusted in existing plants.² The market prices of hydrocarbon

co-products (Price_{*j*}) are based on average European market prices in 2019,¹⁰⁴ with the market price of ethylene itself (Price_{C₂H₄}) at £1000 t_{C₂H₄}⁻¹. The price of O₂ co-product (Price_{O₂}) is taken at £130 t_{O₂}⁻¹.¹⁰⁵ All of the relative prices (Ratio_{*j*}, eqn (3)) are assumed to remain constant throughout the plant's lifetime. Notice, in particular, that ethylene has the highest market price amongst the co-products.

To determine the extent to which the process ranking is subject to key modelling assumptions, a sensitivity analysis is conducted in terms of the LCOE, the stack cost (Cost_{stack}), and the relative market prices of propylene (Ratio_{C₃H₆}) and oxygen (Ratio_{O₂}) with respect to the ethylene price. A Monte Carlo simulation is conducted based on low-discrepancy Sobol sampling with 1024 scenarios, leading to a confidence interval smaller than 1.5% around the mean LCOEt value. All four uncertain factors are modelled with triangular probability distributions (*cf.* Table 2). The costs of uninterrupted renewable electricity (Cost_{el}), green H₂ (Cost_{H₂}), and CO₂ feedstock (Cost_{CO₂}) are assumed to be correlated with the LCOE of offshore wind, whose value ranges between current costs (£59.0 MW h⁻¹) and projections for 2050 (£38.4 MW h⁻¹).^{75,107} Cost_{H₂} is taken directly proportional to Cost_{el} since electricity is the main driver of green H₂ price, while the variation range for Cost_{CO₂} accounts for the fact that electricity contributes around 40% of the cost of the solid sorbent-based DAC process.⁹⁹ Since public data on propylene prices are limited, a range of ±20% is assumed.⁵⁸ The upper and lower bounds for O₂ price are so chosen to describe a scenario whereby the market price would decrease as a result of market saturation.⁴⁹ The stack cost is varied between the capital cost of PEM electrolyser systems in 2020⁷⁵ and projected capital cost of a large-scale plant by 2030.³¹

3 Results and discussion

3.1 Process inventories

Process inventories for the nine selected routes at scale are shown in Fig. 2, with more details about the mass and energy



balances available in Section S6 of the ESI.† All these inventories are scaled for 1 tonne of ethylene produced. Differentiated by colour, the processes correspond to three fully thermocatalytic routes (MS+MTO, CTM+OCM, D-FT), three fully electrocatalytic routes (eCO+S-eC₂H₄, HT-eCO+S-eC₂H₄, D-eC₂H₄), and three hybrid routes (eCO+S-FT, HT-eCO+S-FT, eCH₄+OCM). The input-output inventory streams comprise electricity (subscript el), carbon dioxide (subscript CO₂), hydrogen (subscripts H_{2,f} and H_{2,h}), ethylene (subscript C₂H₄), other hydrocarbons (subscript hc), oxygen (subscript O₂), wastewater (subscript ww), and residual emissions (subscript CO₂-eq).

Since CO₂ provides the carbon feedstock, the CO₂ inventories closely correlate with the total hydrocarbon production (sum of C₂H₄ and hc). Most of the unconverted CO₂ and intermediates are recycled within the process boundaries, therefore the net CO₂ feedstock requirement is largely independent of the overall conversion rates. Despite the reference processes for D-FT and S-FT being tailored towards light olefins, their ethylene yields remain low and large amounts of CO₂ are consumed in side reactions to co-products such as propylene and C₄₊. As a result, D-FT, eCO+S-FT and HT-eCO+S-FT have higher CO₂ feedstock requirements than other routes. Likewise, the MS+MTO route has a large CO₂ requirement due to significant propylene and C₄₊ side-production. Conversely, the electrocatalytic processes eCO+S-eC₂H₄, HT-eCO+S-eC₂H₄ and D-eC₂H₄ require less CO₂ feedstock since ethylene is the single product. Even though ethylene is also the sole product of OCM, the CO₂ requirement of the CTM+OCM and eCH₄+OCM routes is intermediate as part of the unconverted methane is combusted in a turbine for electricity production, thus venting CO₂ with the exhaust gas.

Hydrogen is either consumed as reaction feedstock (H_{2,f}) or used as heating utility (H_{2,h}). Neither of the electrocatalytic processes (eCO+S-eC₂H₄, HT-eCO+S-eC₂H₄, D-eC₂H₄) nor the methane-mediated route (eCH₄+OCM) require hydrogen as feedstock since hydrogen is generated *in situ* from the humidified CO₂ feed. However, as all low-temperature electrocatalytic processes have low single-pass conversion rates, they require hydrogen to provide heat for the regeneration of the amine sorbent used in the separation of unreacted CO₂ before recycling. By contrast, the thermocatalytic processes consume hydrogen mainly as a feedstock. Given that the reactions are typically exothermic, the hydrogen feedstock is also indirectly used to meet part of the heat and electricity demands through process heat integration. However, additional hydrogen as heat utility is required for the D-FT process to preheat its large recycle stream and for the MS+MTO process despite using excess heat to preheat the reactor feed (*cf.* Section S6 of the ESI†). All of the thermocatalytic processes furthermore include turbines for converting excess reaction heat into electricity. But only for the integrated CTM+OCM process is the excess heat high enough for an electricity surplus to be generated and sold to the grid. The consumption pattern of the hybrid process eCO+S-FT reflects the characteristics of thermo- and electrocatalytic routes because significant amounts of hydrogen are consumed both as feedstock to achieve the necessary syngas

composition and as heating fuel in the carbon capture process due to the low single-pass conversion of the eCO stage.

The electricity demands of the electrocatalytic and hybrid processes correlate with their cell voltages and faradaic efficiencies. Despite the high faradaic efficiency of the eCO and HT-eCO stages and the excess electricity generated in the S-FT stage (15.9 MW h_{el} per t_{CO}), eCO+S-FT and HT-eCO+S-FT have the highest electricity demand due to their large intermediate syngas stream (33.6 t_{CO} per t_{C₂H₄}); eCH₄+OCM comes in third position as the eCH₄ stage has by far the highest cell voltage. In contrast, the fully electrocatalytic routes eCO+S-eC₂H₄, HT-eCO+S-eC₂H₄ and D-eC₂H₄ consume 3 to 4 times less electricity than the hybrid routes. And although eCO+S-eC₂H₄, HT-eCO+S-eC₂H₄ require 10e⁻ per mol_{C₂H₄} compared to 12e⁻ per mol_{C₂H₄} in D-eC₂H₄, all three have similar electricity demands. For those processes having electrocatalytic conversion stages, the quantity of the oxygen by-product is commensurate to the electricity demand given the OER is the complementary half-cell reaction of the CO₂/CO reduction reaction.

A significant amount of water is formed as by-product in both thermocatalytic and electrocatalytic processes. These wastewater streams either contain traces of hydrocarbons and alcohols in the case of thermocatalytic processes or correspond to an alkaline solution in the case of electrocatalytic processes, and they require treatment before crossing the process boundaries. Although the costs for an external wastewater treatment service were included in the TEA, on-site wastewater regeneration and recycling could also be implemented to lower the water footprint of the processes. The amount of wastewater produced by thermocatalytic processes generally correlates with the amount of hydrocarbons formed. The CTM+OCM process uses large amounts of steam to dilute the OCM reactor feed and therefore generates more wastewater than the MS+MTO process. Recall also that the wastewater stream of the D-FT process could not be quantified due to incomplete mass balances (*cf.* Section A.4). But as dewatering at the S-FT reactor outlet alone produces 18.81 t_{H₂O} per t_{C₂H₄}, wastewater production is expected to be comparable for D-FT.

Direct (gross) emissions from the thermocatalytic processes are caused by purge streams, which are translated into CO₂-equivalents on Fig. 2; while in electrocatalytic processes, the direct emissions correspond to any residual CO₂ not captured by the mixed-amine absorption column. These direct emissions are lower than those of conventional ethane crackers (c. 0.74 t_{CO₂} t_{C₂H₄}⁻¹)¹⁰⁸ for both the direct electrochemical route D-eC₂H₄ and the thermochemical route MS+MTO, and slightly higher for the two-stage electrochemical routes eCO+S-eC₂H₄ and HT-eCO+S-eC₂H₄. By contrast, the hybrid processes eCO+S-FT and HT-eCO+S-FT present much larger direct emissions due to the Fischer-Tropsch synthesis having a small ethylene yield and forming a range of saleable co-products. Nevertheless, an economic allocation among these co-products (*cf.* Fig. 4 below) suggests that the direct emissions associated with ethylene would remain comparable to that of ethane cracking. Recall also that the purge stream and hence the emissions of the D-FT process could not be quantified due to incomplete mass



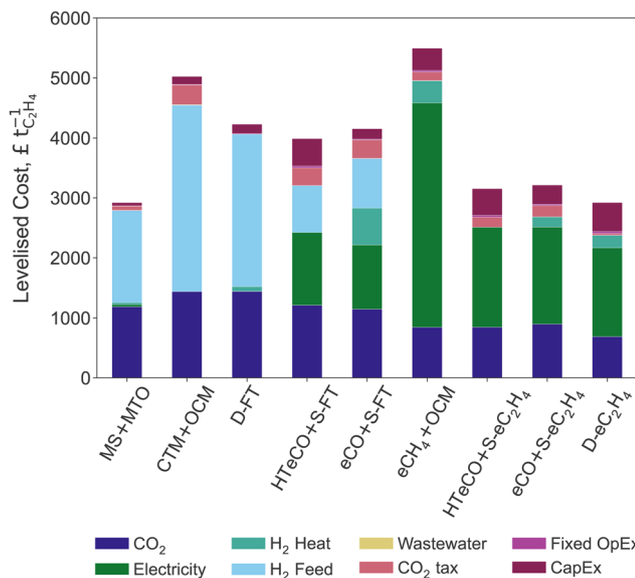


Fig. 3 Levelised cost of ethylene and corresponding breakdown for the nine candidate routes.

balances (*cf.* Section A.4). Lastly, the carbon intensity of both the CTM+OCM and eCH₄+OCM processes exceeds that of ethane cracking because significant CO₂ emissions are caused by the combustion of unconverted methane from the OCM stage to generate electricity. Although not implemented here, capturing and recycling CO₂ from this flue gas might be economically competitive given the high cost of DAC. It is also worth reiterating that the CO₂ feedstock is much larger than the direct (gross) emissions in all of the green ethylene routes and so, unlike ethane crackers, the production processes are still net CO₂ consumers. To draw final conclusions on the environmental impact, a thorough life-cycle assessment should be performed in future research.

3.2 Techno-economic comparison

The levelised costs of ethylene (LCOEt) from DAC and renewable electricity for the nine production pathways are compared in Fig. 3. The breakdown differentiates capital, fixed and variable operating costs, the latter being further differentiated into the contributions of CO₂ feedstock, renewable energy (electricity, hydrogen), wastewater treatment, and carbon tax.

The predicted LCOEt ranges between £2920–5280 t_{C₂H₄}⁻¹. Remarkably, both the thermocatalytic process MS+MTO and the direct electroreduction process D-eC₂H₄ provide the lowest-cost options at around £2920 t_{C₂H₄}⁻¹. The most expensive routes are the methane-mediated hybrids with OCM, eCH₄+OCM and CTM+OCM, both above £5000 t_{C₂H₄}⁻¹. The Fischer–Tropsch routes D-FT, eCO+S-FT and HT-eCO+S-FT have a comparable LCOEt in the range of £3930–4230 t_{C₂H₄}⁻¹. Given that the LCOEt of D-FT omits the carbon tax for direct emissions and wastewater treatment cost, its LCOEt is however underestimated. The three electrocatalytic routes D-eC₂H₄, eCO+S-eC₂H₄ and HT-eCO+S-eC₂H₄ also perform similarly with an LCOEt between £2920–3220 t_{C₂H₄}⁻¹.

Table 3 Capital expenditures (CapEx) for the nine candidate routes at a production capacity of 800 kt year⁻¹

Route	CapEx, £ ₂₀₁₉ Bn
MS+MTO	1.17
CTM+OCM	1.21
D-FT	5.72
HT-eCO+S-FT	37.81
eCO+S-FT	13.56
eCH ₄ +OCM	7.80
HT-eCO+S-eC ₂ H ₄	6.54
eCO+S-eC ₂ H ₄	4.88
D-eC ₂ H ₄	7.54

All of these levelised costs can be interpreted as lower bounds on green ethylene market prices since they omit any profit margins. Even compared to a high market price of £1400 t_{C₂H₄}⁻¹ (recorded in April 2022)¹⁰⁹ and after taxing the CO₂ emissions of a conventional ethane cracker (c. £300 t_{C₂H₄}⁻¹ at 0.97 t_{CO₂} t_{C₂H₄}⁻¹), the predicted green ethylene prices are far from being competitive with fossil-based ethylene.

The routes involving an electrocatalytic conversion step generally have a higher CapEx than the thermocatalytic processes (see Table 3). The eCO+S-eC₂H₄ process achieves the lowest CapEx among the electrocatalytic and hybrid routes because it requires a relatively small electrode area given its high current density (*cf.* Section 2.1.2). It is nevertheless 4 times more expensive than MS+MTO or CTM+OCM. Despite being fully thermocatalytic, D-FT requires larger unit and equipment sizes to make up for its low ethylene yield, which is why its CapEx is comparable to that of eCO+S-eC₂H₄ and 3.5 times greater to that of S-FT. The CapEx of the electrocatalytic and hybrid processes are dominated by the CO₂/CO electrolyser cost, while the separation equipment and thermocatalytic stage contribute less than 6% and 12%, respectively.

Despite the large differences observed in Table 3, the annualised CapEx only represents a minor share of the LCOEt for any of the processes (see breakdown in Fig. 3). The contributions of the fixed OpEx are also minor as they correspond to a small percentage of the CapEx (*cf.* Section 2.2). This conclusion holds insofar as the manufacturing of CO₂/CO electrolysers can benefit from the learning rates that are predicted for PEM and SOEC electrolysers. It is also worth recalling that the assumed 5-year lifetime of the electrolyser stack by 2035 (*cf.* Section S2 of the ESI[†]) far exceeds current stability reports for low-temperature CO₂/CO electrolyser cells.^{71,72,110,111} For instance, reducing the life-time of the D-eC₂H₄ electrolyser stack to 4 years is predicted to increase its CapEx by 19%. Therefore, extending the lifetime of the electrolyser stack is key towards improving the cost-competitiveness of electrocatalytic routes.¹¹²

It is clear from Fig. 3 that the cost of renewable energy—both renewable electricity and green H₂—makes up the largest share of the LCOEt across all nine routes. For the fully thermocatalytic routes (MS+MTO, CTM+OCM, D-FT), green H₂ contributes over 50% of the LCOEt. Instead, renewable electricity contributes about 50% of the LCOEt for the fully electrocatalytic routes (eCO+S-eC₂H₄, HT-eCO+S-eC₂H₄, D-eC₂H₄). The LCOEt of the hybrid route eCH₄+OCM is also dominated by the



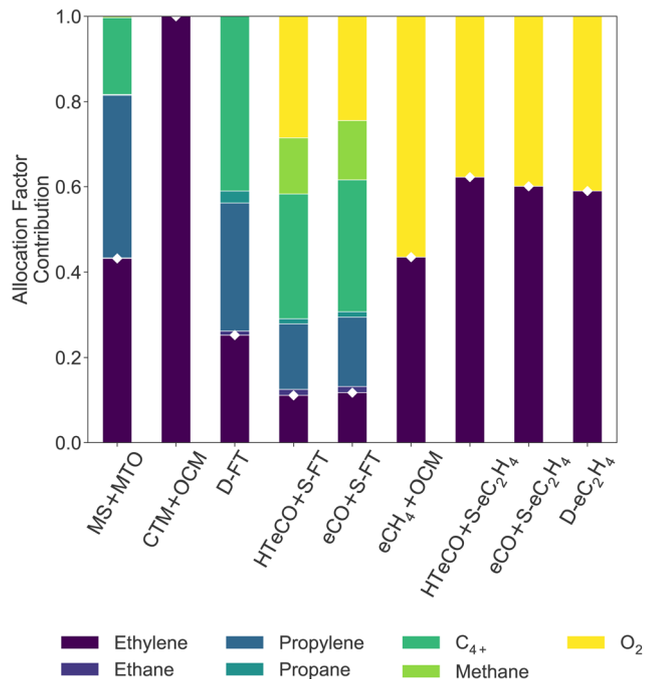


Fig. 4 Economic allocation factors AF_j (eqn (2)) of the saleable co-products in all nine candidate routes. The allocation factor of ethylene ($AF_{C_2H_4}$, white diamonds) determines the LCOEt (eqn (1)).

electrochemical stage eCH_4 due to its high cell voltage (*cf.* Table 1), while the procurement of hydrogen and electricity both contribute significantly to the LCOEt of $eCO+S-FT$ and $HT-eCO+S-FT$. Except for $eCO+S-FT$ (15%), the contribution of hydrogen as a heating utility to the LCOEt remains below 7%.

In all nine candidate routes, the CO_2 feedstock has a notable contribution on the LCOEt because of the high cost of DAC. Substituting CO_2 from DAC for fossil-based CO_2 could reduce the share of CO_2 feedstock to 3–10% of the LCOEt, thereby making its contribution significantly smaller than the cost of green electricity or hydrogen. Conversely, the residual emissions make a minor contribution to the LCOEt in any of the selected routes, not only for a carbon tax of $\pounds 301.4 \text{ t}_{CO_2}^{-1}$ (Table 2) but also in the high price scenario of BEIS with $\pounds 504 \text{ t}_{CO_2}^{-1}$ by 2035.¹⁰² Making MS+MTO cost-competitive with ethylene from a conventional ethane cracker (c. $0.74 \text{ t}_{CO_2} \text{ t}_{C_2H_4}^{-1}$) would entail a carbon price of $\pounds 640 \text{ t}_{CO_2}^{-1}$ or higher.

While the processes differ in conversion and ethylene yield, the LCOEt is ultimately governed by its economic allocation factor ($AF_{C_2H_4}$, eqn (4)), which reflects the relative market prices of the co-products. A breakdown of the economic allocation factor for all nine routes is shown on Fig. 4. CTM+OCM is the only selected route with ethylene as the single saleable product ($AF_{C_2H_4} = 100\%$). All the electrocatalytic routes produce oxygen as a saleable by-product alongside the desired carbon monoxide or hydrocarbons, leading to $AF_{C_2H_4}$ values between 45–65%. Propylene and C_4+ are the dominant co-products of the thermochemical routes MS+MTO and D-FT as well as the hybrid route $eCO+S-FT$. While MS+MTO has an $AF_{C_2H_4}$ value over 40%, those of the Fischer-Tropsch syntheses are much lower due to low ethylene yields.

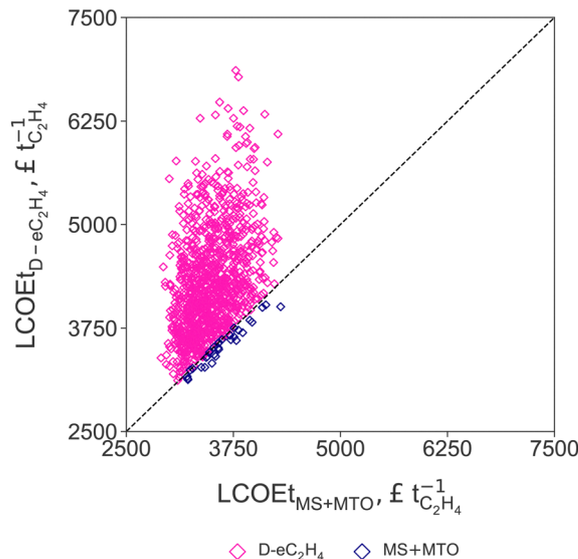


Fig. 5 Parity plot comparing the levelised cost of ethylene (LCOEt) for the thermocatalytic production of ethylene via methanol (MS+MTO, on x-axis) and direct CO_2 electroreduction to ethylene (D- eC_2H_4 , on y-axis) for each uncertainty scenario. The levelised cost of electricity (LCOE), AEM stack cost, and market prices of propylene and oxygen are varied simultaneously (*cf.* Table 2).

Based on this techno-economic comparison, it is unlikely for green ethylene production involving a Fischer-Tropsch synthesis (D-FT, $eCO+S-FT$, $HT-eCO+S-FT$) to be cost-competitive with other thermocatalytic processes, unless the value of its co-products or its conversion rate were to increase significantly. Neither of the OCM routes (CTM+OCM, eCH_4+OCM) is furthermore competitive with MS+MTO given that OCM does not produce any other valuable hydrocarbon products than ethylene. The cost advantage of MS+MTO over CTM+OCM, eCH_4+OCM or D-FT is consistent with previous techno-economic assessments.^{44,45}

Finally, the LCOEt of all three electrocatalytic routes ($eCO+S-eC_2H_4$, $HT-eCO+S-eC_2H_4$, D- eC_2H_4) is within 10% of that of MS+MTO. This gap would increase significantly if the oxygen by-product from the electrolysers could no longer be valorised or if the life-time of the electrolyser stack did not extend beyond 3 or 4 years, as already noted. A single-pass conversion lower than the assumed 40% in AEM electrolysers (*cf.* Table 1) would also be detrimental to the LCOEt, due to the combined effect of increasing H_2 consumption to recycle unconverted CO_2 , CO_2 feed and CO_2 emissions. For instance, a 30% single-pass conversion would already increase the LCOEt of both D- eC_2H_4 and $eCO+S-eC_2H_4$ by 5%. The overall similarity in LCOEt between $eCO+S-eC_2H_4$, $HT-eCO+S-eC_2H_4$ and D- eC_2H_4 is in agreement with recent techno-economic assessments, albeit conducted under different system boundaries.^{48,113} If it were not for the high electrolyte molarity (7 M KOH) in the selected D- eC_2H_4 reference (*cf.* Section A.4 of the Appendix),⁷² D- eC_2H_4 might as well rank behind the two tandem routes $eCO+S-eC_2H_4$ and $HT-eCO+S-eC_2H_4$. For instance, in a scenario with 1 M KOH electrolyte and target cell voltage of 3.5 V,^{114,115}



the LCOEt of D-eC₂H₄ would already increase by 17% (£3600 t_{C₂H₄}⁻¹).

3.3 Sensitivity analysis

Results from the Monte Carlo analysis results for the two routes with the lowest LCOEt (MS+MTO and D-eC₂H₄) are summarised in Fig. 5, where uncertainties in the LCOE, stack cost, and market prices of propylene and oxygen relative to the ethylene price are varied simultaneously (*cf.* Section 2.3). The parity plot confirms that the production cost of the MS+MTO process remains the lowest across the range of sampled uncertainty scenarios. Only in scenarios with low propylene price, low stack cost, high LCOE and high oxygen price, does the LCOEt of D-eC₂H₄ become comparable to that of MS+MTO. This scenario analysis also reveals that the production cost of green ethylene at three times the current market price of (fossil-based) ethylene corresponds to a rather favourable scenario, due in large part to the skewed distributions of LCOE, stack cost and oxygen price.

Recall that varying the LCOE of offshore wind between current costs (£59.0 MW h⁻¹, +40%) and projections for 2050 (£38.4 MW h⁻¹, -9%) changes the cost of non-intermittent renewable electricity, green hydrogen and air-captured CO₂ proportionally (*cf.* Section 2.3). The LCOEt of both MS+MTO and D-eC₂H₄ processes correlate linearly with the LCOE (Fig. 6A). In particular, the ranking between MS+MTO and D-eC₂H₄ is preserved over the whole LCOE range. D-eC₂H₄ could become the cheapest option if the faradaic efficiency were to reach 80% (from the nominal 66% efficiency). But if PEMs were used instead of AEMs, the cell potential would increase by 25%, and the LCOEt gap between the two processes would widen; as it would if the electrolyte molarity was reduced to 1 M KOH (from the nominal 7 M KOH, *cf.* Section 3.2). Nevertheless, a more detailed modelling would be required to quantify the potential benefits afforded by adapting the operation of D-eC₂H₄ to fluctuating electricity prices.

Reducing the stack cost from current prices to costs projected for 2030 would help close the LCOEt gap, but still not enough for D-eC₂H₄ to overtake MS+MTO (Fig. 6B). For D-eC₂H₄ to gain a competitive advantage over MS+MTO, reductions by over 12% in electricity consumption would be necessary, through improvement in faradaic efficiency and cell voltage. Given that the annualised CapEx contributes only 17% to the LCOEt (Fig. 3), the LCOEt of D-eC₂H₄ is however more sensitive to variations in the relative market price of oxygen than it is to the stack cost (compare Fig. 6B and D).

The large amount of oxygen by-product (5.2 t_{O₂} t_{C₂H₄}⁻¹) makes its market price decisive for the economic allocation factor of D-eC₂H₄ (*cf.* Section 3.2). If the oxygen price were to drop as a consequence of market saturation, the LCOEt of D-eC₂H₄ would increase, even causing the LCOEt to double in scenarios where the oxygen price is null (Fig. 6C). Similarly, if the market price of oxygen were to remain at today's level (£130 t_{O₂}⁻¹) while the market price of green ethylene tripled (£3000 t_{C₂H₄}⁻¹), the LCOEt of D-eC₂H₄ would be double that of MS+MTO.

Propylene is the main saleable co-product of the MS+MTO route and thereby influences the economic allocation factor. A negative correlation between the price ratio and the LCOEt is observed in Fig. 6D. But with about 1 t_{C₃H₆} produced for each t_{C₂H₄}, propylene price variation exerts a smaller effect on MS+MTO than the oxygen price does on D-eC₂H₄ (compare Fig. 6C). Even for a propylene price on par with ethylene, as observed for instance in April 2022,¹¹⁶ the LCOEt from MS+MTO would only decrease by a mere 2.3%.

Our finding that MS+MTO remains the lowest-cost process for green ethylene production across the majority of simulated scenarios suggests that scaling up of the current best available CO₂ electrolyzers may not be enough for producing low-cost green ethylene and significant performance improvement is needed for electrocatalytic routes to become competitive with MS+MTO. National green hydrogen production targets carry an additional risk to the competitiveness of D-eC₂H₄ as oxygen is also a by-product of water electrolysis. If the UK's ambitious target of 10 GW of low-carbon hydrogen production were to be produced in the form of green hydrogen, an extra 2 Mt_{O₂} per year would be added to the market by 2030. Without a concurrent increase in demand, *e.g.* for oxy-combustion¹¹⁷ or in wastewater treatment applications,¹¹⁸ the oxygen market price could drop. Meanwhile, the prices of green ethylene produced by MS+MTO could drop if the propylene price were to rise. Since ethylene and propylene are predominantly co-produced in naphtha crackers, their prices in the European market are currently highly correlated. But this interdependence could weaken in the future if the defossilisation of ethylene and propylene relied on separate feedstock. As indicated by the cost breakdown in Fig. 3 though, the LCOE from renewable power generators remains the most sensitive factor for competitive green ethylene prices by and large.

3.4 Beyond the process boundaries

At the time this paper is written, all three ethane steam crackers in the UK are reliant on shale gas imports from the US.^{119,120} CO₂ utilisation for ethylene production therefore would promote the use of local resources such as offshore wind power. But a major impediment to the deployment of green ethylene lies with the huge demand for renewable electricity. The modelling conducted in the previous sections predicts that MS+MTO and D-eC₂H₄ would consume, respectively, 52 TW h_{el} and 41 TW h_{el}—assuming an efficiency of 1.3 MW h_{el} MW h_{H₂,HHV}⁻¹ for PEM electrolyzers—to displace a single cracker with an annual capacity of 800 kt. In particular, these demands correspond to 13–16% of the UK's total electricity generation in 2022 (325 TW h_{el}).¹²¹ On top of this, powering the DAC plant for MS+MTO and D-eC₂H₄ would require an additional 14 TW h_{el} and 6 TW h_{el} of renewable electricity, respectively, even with efficient heat pump systems and assuming an average electricity requirement of 2 MW h_{el} t_{CO₂}⁻¹.⁹⁹ These figures also need to be contrasted with our reference ethane cracker in Wilton (UK) consuming 11.5 TW h_{el} per year in total,⁶ a much smaller share (*c.* 3.5%) of the UK's annual electricity consumption.



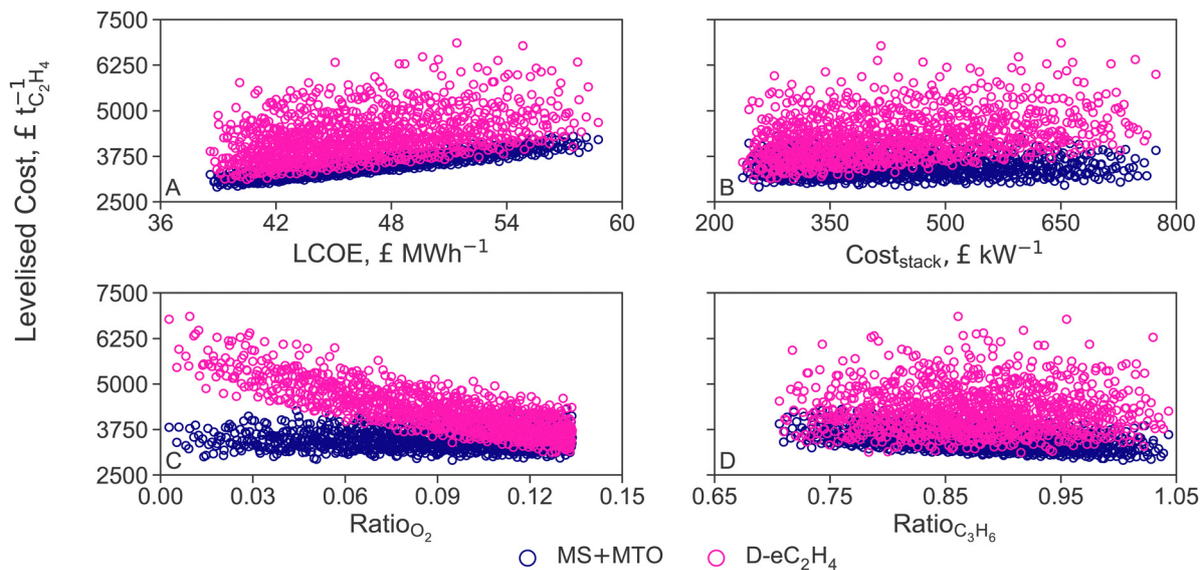


Fig. 6 Sensitivity analysis of the levelised cost of ethylene (LCOEt) to variations in the levelised cost of electricity (LCOE, A), AEM stack cost (B) and market prices of oxygen (C) and propylene (D) for the thermocatalytic production of ethylene *via* methanol (MS+MTO) and direct CO₂ electroreduction to ethylene (D-eC₂H₄). The costs of hydrogen and CO₂ feedstock are varied simultaneously with LCOE (*cf.* Table 2).

Alongside the huge demand for renewable energy to power green ethylene comes the physical footprint of a dedicated offshore wind farm. Assuming an average power per unit area of 3 W m⁻² in the UK,¹²² the required offshore area would be 2800 km² for MS+MTO and 2000 km² for D-eC₂H₄, corresponding to 1.2–1.7 times the area of Greater London.¹²³ Such figures put a clear pressure on the chemical industry to take a strategic decision on securing its supply of low-cost, low-carbon electricity. Delaying this decision could put the industry at risk of missing out on the best offshore wind farm leases, as the seabed areas in the UK exclusive economic zone (EEZ) are continuously auctioned. In the context of low-carbon ethylene production, the chemical industry might want to reconsider the idea of co-locating future chemical plants with nuclear plants that could supply both heat and power.¹²⁴

Both geographic and geological characteristics govern the cost of green ethylene production. Like the UK, Northern Germany and Denmark have large offshore wind potentials in close proximity to underground salt caverns for low-cost, large-capacity hydrogen storage.¹²⁵ Therefore, a LCOEt similar to our case study (Fig. 3) could be expected for production sites in those countries. While salt caverns are also available offshore in the North Sea, higher cost for offshore solution mining, injection wells and pipelines would exacerbate the cost of hydrogen storage. For production sites where geological hydrogen storage is unavailable, the use of pressure tanks could increase the hydrogen cost by 20%,⁹⁸ in turn increasing the LCOEt of MS+MTO by 10%.

At the global level, the production cost of green ethylene will be dependent on the availability of renewables, climatic conditions, and financing conditions all together. Outside Europe, the Western US might offer suitable conditions for a similar LCOEt as in the UK (Fig. 3), but given that fossil-based ethylene in the US is less than half the price on the European market,¹²⁶

green ethylene would be even less competitive on the US market. Comparable LCOE and LCOH to the UK could also be achieved in other parts of the World, such as the West Coast of Canada, the Arabic Peninsula or Australia.^{127,128} Nevertheless, the climatic conditions there would make CO₂ capture using solid sorbent-based DAC 10–25% more expensive.⁹⁹ A more detailed and systematic assessment is necessary to identify favourable locations for green ethylene production.

When deciding between D-eC₂H₄ and MS+MTO for the production of green ethylene, not only the cost but also the supply security during dark doldrums, periods in which little to no electricity is generated by offshore wind farms, should be considered. Renewable electricity is a localised resource that may not be transported over long-distances and for which supply and demand need to be matched at any point in time. Instead, local production of green hydrogen could be supplemented by imports from countries with similar or lower predicted production costs, such as Algeria and Australia.^{128,129} Even though hydrogen imports could also be reconverted to electricity for D-eC₂H₄, the associated losses would make it more cost-effective to directly use hydrogen in the MS+MTO process. For a future in which the chemical industry is mainly reliant on renewable power, a higher diversity of supply could bring greater price stability, as the price of commodity chemicals such as ethylene are otherwise at risk of becoming weather-dependent.

Finally, the main motivation behind displacing fossil-based ethylene for green ethylene lies in the prospect of producing carbon-negative ethylene and co-products (use phase excluded). To calculate the net CO₂ emissions, the CO₂ inlet streams are subtracted from the sum of scope-1 residual emissions and scope-2 emissions associated with electricity that is consumed either directly or for the production of H₂ and CO₂. Assuming an average carbon intensity of 7.5 kg_{CO₂-eq} MW h_{el}⁻¹,¹³⁰ the



allocated net emissions per tonne of ethylene produced amount to $-3.3 t_{\text{CO}_2\text{-eq}}$ for MS+MTO and $-1.9 t_{\text{CO}_2\text{-eq}}$ for D-eC₂H₄. Both thermocatalytic and electrocatalytic routes would thus present a negative carbon balance from a cradle-to-gate point of view. On the path to net-zero, this would allow for a dramatic reduction over conventional ethane steam crackers, whose cradle-to-gate emissions are estimated to exceed $3 t_{\text{CO}_2\text{-eq}}$ per tonne of ethylene with ethane from shale gas.¹⁰⁸ A cradle-to-grave assessment encompassing the ethylene use phase in consumer products and any downstream emissions is nevertheless required to conclude on the negative emission potential of green ethylene as a whole.¹³¹

4 Conclusions

A techno-economic comparison between mature thermocatalytic and novel electrocatalytic processes was conducted for the production of green ethylene from air-captured CO₂ and offshore wind electricity, with a view to displacing a conventional ethane cracker in the UK at the horizon of 2035. Similar to fossil-based processes today, the production cost of green ethylene would be dominated by the cost of energy and carbon feedstock, regardless of the process alternative. The fully-thermocatalytic route converting CO₂ to methanol then to olefins would achieve the lowest cost ($\text{£}2900 t_{\text{C}_2\text{H}_4}^{-1}$), followed by fully-electrocatalytic routes ($\text{£}2900\text{--}3200 t_{\text{C}_2\text{H}_4}^{-1}$). Process routes relying on Fischer–Tropsch synthesis or oxidative coupling of methane would furthermore be non-competitive. Although considering a UK case study herein, similar levelised cost of green ethylene are expected in Northern Germany or Denmark.

Apart from the levelised cost of electricity, the market price of the oxygen by-product and electrolyser stack cost are identified as key sensitive factors to the cost-competitiveness of the electrocatalytic processes. However, the methanol-mediated thermocatalytic route is predicted to out-compete the direct electroreduction of CO₂ to ethylene, even for scenarios of high oxygen prices and low stack cost. From a cradle-to-gate perspective, both processes have the potential to reduce the carbon emissions of ethylene production significantly and achieve net-negative emissions between -3.3 to $-1.9 t_{\text{CO}_2\text{-eq}}$. But a major barrier to green ethylene deployment lies in its energy-intense nature, making it highly reliant on a large-scale supply of low-cost renewable electricity. Displacing a single cracker with an annual production capacity of $800 \text{ kt}_{\text{C}_2\text{H}_4}$ would consume the equivalent of 14–20% of the UK's current power generation and require an offshore wind farm larger than the area of Greater London. Finally, transitioning from fossil fuels to intermittent renewable electricity requires a rethinking of the supply security of chemical processes, especially during dark doldrums. The process inventories developed herein could feed into decision-support models for analysing the wider consequences of ethylene defossilisation for the UK's future energy system. As an interim solution, ethylene production will continue to rely on fossil feedstock but could be retrofitted with CCS or H₂ as fuel to lower its emission intensity.

Nomenclature

Acronyms

CO ₂ RR	Electroreduction of CO ₂
CORR	Electroreduction of CO
AEM	Alkaline electrolyte membrane
AF	Allocation factor
UF	Utilisation factor
ASU	Air separation unit
BEIS	Department for Business, Energy & Industrial Strategy
BOP	Balance of plant
CapEx	Capital expenditures
CCS	Carbon capture and storage
CEPCI	Chemical engineering plant cost index
CRF	Capital recovery factor
DAC	Direct air capture
EEZ	Exclusive economic zone
FT	Fischer–Tropsch
FE	Faradaic efficiency
GHG	Greenhouse gas
HHV	Higher heating value
LCA	Life-cycle assessment
LCOE	Levelized cost of electricity
LCOEt	Levelized cost of ethylene
LCOH	Levelized cost of hydrogen
OER	Oxygen evolution reaction
OpEx	Operational expenditures
PEM	Proton exchange membrane
PTFE	Polytetrafluoroethylene
SOEC	Solid oxide electrolysis cell
TIC	Total invested capital
TRL	Technology readiness level
UK	United Kingdom
US	United States
VPSA	Vacuum pressure swing adsorption

Subscripts

el	Electricity
eq	Equivalents
f	Feedstock
h	Heating
hc	Hydrocarbons other than ethylene
stack	Electrolyser stack
tax	Carbon tax
ww	Wastewater

Model parameters and variables

η_F	Faradaic efficiency
LT	Lifetime
E_{cell}	Cell voltage
E_{red}	Reduction potential
$f_{\text{O\&M}}$	Share of fixed operating costs on CapEx
j	Current density
m_j	Production rate of product j
r	Interest rate
X	Conversion
AF_j	Allocation factor of co-product j



A_{cell}	Area of the electrode
Cost_j	Unit cost of process input/emission j
Price_j	Market price of co-product j
Ratio_j	Ratio of market price of co-product j to ethylene

Process technologies

CTM	CO ₂ methanation
D-eC ₂ H ₄	Direct electroreduction of CO ₂ to ethylene
D-FT	Direct Fischer–Tropsch synthesis
eCH ₃ OH	Electroreduction of CO ₂ to methanol
eCH ₄	Electroreduction of CO ₂ to methane
eCO	Electroreduction of CO ₂ to CO
HT-eCO	High-temperature electroreduction of CO ₂ to CO
MS	Methanol synthesis
MTO	Methanol to olefins
OCM	Oxidative coupling of methane
S-eC ₂ H ₄	Electroreduction of CO to ethylene
S-FT	Syngas Fischer–Tropsch synthesis

Author contributions

Andreas H. Nyhus: data curation, formal analysis, methodology, validation, software, investigation, visualisation, writing – original draft preparation. Maria Yliruka: conceptualisation, supervision, data curation, methodology, investigation, validation, writing – original draft preparation, writing – reviewing and editing. Nilay Shah: funding acquisition, writing – reviewing and editing. Benoît Chachuat: supervision, validation, writing-reviewing and editing.

Data availability

Data is available upon request.

Conflicts of interest

There are no conflicts to declare.

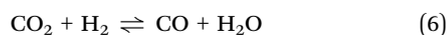
Appendix

A Green ethylene production routes

This appendix provides further details about the selected reference processes in the conversion of CO₂ to green ethylene (*cf.* Fig. 1). TRLs of these processes are summarised in Table 4.

A.1 Methanol intermediate route

Methanol synthesis (MS). Production of methanol *via* CO₂ hydrogenation (reactions (5) and (6)) has reached TRL 9, with commercial plants already operating in China.¹³² CO₂ and H₂ flow over a Cu/ZnO/Al₂O₃ catalyst in a fixed bed reactor under 76 bar and 210 °C to form methanol at a purity of 99.9 wt%.⁹³



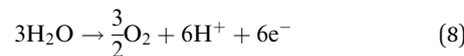
The CO₂ hydrogenation to methanol process was modelled by Matzen *et al.*,¹³³ Wiesberg *et al.*,¹³⁴ and Pérez-Fortes

Table 4 Technology readiness level (TRL) of the process candidates. Refer to Fig. 1 for a list of abbreviations

Process	TRL
MS	9
eCH ₃ OH	4
MTO	9
CTM	9
eCH ₄	4
OCM	6
eCO	4
HT-eCO	8
S-FT	4
S-eC ₂ H ₄	4
D-FT	4
D-eC ₂ H ₄	4

*et al.*⁹³ The first was discarded as the levelised capital cost (£65.1 t⁻¹) was significantly larger than those in Wiesberg *et al.*¹³⁴ (£20.5 t⁻¹) and Pérez-Fortes *et al.*⁹³ (£19.3 t⁻¹). Since Wiesberg *et al.*¹³⁴ did not integrate the waste heat of the reactor, the process model from Pérez-Fortes *et al.*⁹³ minimising exergy losses was selected as reference.

CO₂RR to methanol (eCH₃OH). Electroreduction of CO₂ to methanol (half-cell reactions (7) and (8)) over a Cu catalyst¹³⁵ is most effective with a proton-exchange membrane (PEM) and alkaline electrolyte in a flowing cell set-up (TRL 4).^{136–139}



In contrast to other electrocatalytically produced intermediates, methanol is found in the electrolyte together with water and formic acid as common side-products.¹³⁷ Based on the low single-pass production rate ($3.2 \times 10^5 \text{ mol m}^{-2} \text{ s}^{-1}$) reported in Albo *et al.*,¹⁴⁰ the electrolyte would need too large a recirculation ratio in order to increase methanol concentration before distillation. This route was therefore excluded from further analysis on the basis of technical infeasibility.

Methanol to olefins (MTO). Production of light olefins from methanol is at TRL 9, with commercial plants in operation in China.¹⁴¹ A SAPO-34 catalyst¹⁴² enables the conversion of methanol and steam at 3 bar and 400 °C to light olefins, achieving close to 100% conversion and 80% selectivity to ethylene and propylene at a product purity of 99.5 wt%.¹⁴³ The hydrocarbon products [CH₂] are formed *via* the dimethylether intermediate CH₃OCH₃ according to a complex exothermic reaction scheme,¹⁴² as summarised in reaction (9).



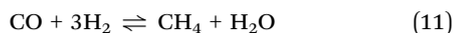
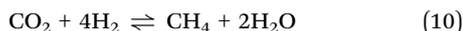
Economic analyses of the MTO process are found in Jasper *et al.*¹⁴⁴ and Chen *et al.*¹⁴³ The first was discarded as the authors significantly simplified the actual plant by adjusting stream compositions using hydrocarbon imports and simplifying the separation train to three columns instead of five.¹⁴² Of the two designs presented in Chen *et al.*,¹⁴³ the optimised



design was chosen as reference due to its much reduced energy requirement of 0.56 GJ t^{-1} compared to 1.51 GJ t^{-1} in the second one.

A.2 Methane intermediate route

CO₂ methanation (CTM). Methanation of CO₂ has been commercially operated in Germany since 2013,¹⁴⁵ thus considered TRL 9. Methane is formed by reacting H₂ and CO₂ in a stoichiometric ratio of 4 (reactions (6), (10) and (11)) over a Ni/MgAl₂O₄ catalyst in an adiabatic fixed-bed reactor at 10 bar and at 350 °C.⁹²



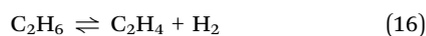
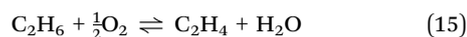
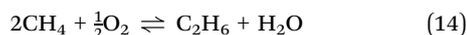
Both Chauvy *et al.*⁹² and Iaquaniello *et al.*¹⁴⁶ modelled CO₂ methanation to produce synthetic natural gas using power-to-gas technology. The latter considered feeding a flue gas stream containing only 8.7 vol% CO₂ and was discarded since the reactor and separation train would be significantly oversized as a result. By contrast, the CO₂ feed stream in Chauvy *et al.*⁹² was 98 mol% pure, so it was selected as reference.

CO₂ electroreduction to methane (eCH₄). Electroreduction of CO₂ to methane (half-cell reactions (12) and (13)) appears as the most immature technology of all the considered CO₂RR routes (TRL 4). Recent research has been more focused on other CO₂RR products such as CO or C₂H₄ than CH₄.¹⁴⁷



For the electrocatalytic conversion of CO₂ to CH₄, Zhang *et al.*⁷⁰ achieved high current densities by coating the Cu/CuO_x catalyst with a hydrophobic carbon shell, while Pan *et al.*¹⁴⁸ achieved low cell voltage and high faradaic efficiency using a Nafion-covered Cu-electrode. The work by Zhang *et al.*⁷⁰ was selected over Pan *et al.*¹⁴⁸ given the reported high faradaic efficiency and a two orders of magnitude higher current density (5000 A m^{-2}). Zhang *et al.*⁷⁰ has the highest current density out of the papers reviewed by Li *et al.*¹⁴⁷

Oxidative coupling of methane (OCM). This process was demonstrated at pilot scale (TRL 6) by Siluria Technologies in the US.¹⁴⁹ Ethylene is produced by reacting CH₄ and O₂ (reactions (14)–(16)) over a Na₂WO₄-Mn/SiO₂(W-Mn/SiO₂) catalyst in a cooled reactor with inlet at 350 °C and outlet at 305 °C under 9 bar.¹⁵⁰

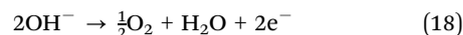


Both Spallina *et al.*¹⁵⁰ and Layritz *et al.*¹⁵¹ modelled OCM as a decarbonisation route for conventional naphtha crackers. The second was discarded because the capital cost of the OCM reactor could not be separated from the total system cost. Of the five process designs compared in Spallina *et al.*,¹⁵⁰ the

reference was chosen as the design optimised for a lower operating temperature.

A.3 Carbon monoxide intermediate route

Low-temperature CO₂ electroreduction to CO (eCO). Common amongst experimental studies on CO₂ electroreduction to CO (half-cell reactions (17) and (18); TRL 4) is the use of a flowing alkaline electrolyte,^{152–154} sometimes paired with an ionic liquid.^{155,156} Candidate transition metal catalysts include Zn,¹⁵³ Ni,¹⁵⁴ Ag,¹⁵⁵ Co,⁶⁹ and Au.¹⁵⁷



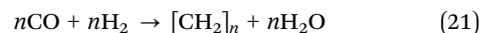
Wang *et al.*⁶⁹ achieved high faradaic efficiency and current density with a CoN₄ catalyst on carbon nanotubes in an alkaline electrolyte membrane (AEM), while Verma *et al.*¹⁵⁷ reported low cell voltage with an Au catalyst on polymer and carbon nanotubes also in an AEM. The former was chosen since the reported current density (3500 A m^{-2}) was 14-times higher.

High-temperature CO₂ electroreduction to CO (HT-eCO). SOEC technology relies heavily on decades-long research and development effort that has gone into commercialising solid oxide fuel cells (SOFC). It has reached TRL 8 for dry electrolysis of CO₂ to CO (half-cell reactions (19) and (20)), for instance with Haldor Topsoe's patented carbon monoxide generator, eCOs™ for small-scale applications.^{83,158} This high-temperature electrolysis operates at 600–850 °C where it benefits from a reduced theoretical minimum cell voltage and faster reaction kinetics.⁸⁰ In contrast to low-temperature CO₂ electrolysis which require a liquid (aqueous) electrolyte, the oxide ions migrate through a ceramic electrolyte in a SOEC, most commonly Y₂O₃-stabilised ZrO₂ (YSZ). The CO₂ electrode needed to conduct both ions and electrons is therefore a ceramic-metal material, usually Ni-YSZ.¹⁵⁹ Often separated by gadolinium-doped cerium oxide layer,¹⁶⁰ a layer of lanthanum strontium manganite (LSM) perovskite can serve as the anode.¹⁶¹



The selected SOEC model relies on public data reported by Haldor Topsoe,¹⁵⁸ specifically the upper bound on the total energy consumption (8 kW h per Nm^3 of CO produced) in order to achieve a high-purity CO stream (>99% purity).

Syngas Fischer–Tropsch synthesis (S-FT). While FT synthesis is already used at commercial scale for fuel production, its application in olefins production is still at the research stage (TRL 4).^{162,163} The exothermic conversion of syngas to low olefins (reaction (21)) over a Fe–Mn–Cu–K catalyst has been reported under 10 bar and 330 °C.⁵⁷

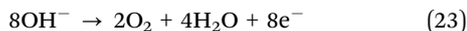
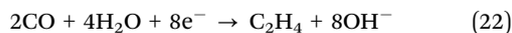


Liu *et al.*⁵⁷ and Xiang *et al.*¹⁶⁴ both studied Fischer–Tropsch synthesis of olefins, including the generation of syngas from fossil-fuels within the system boundary. The former was



selected as reference since the natural gas reforming step could be excluded from the rest of the process.

CO electroreduction to ethylene (S-eC₂H₄). Cu has emerged as the catalyst of choice for CO reduction to ethylene under alkaline conditions (half-cell reactions (22) and (23)) in lab-scale electrolyzers (TRL 4).^{71,165–167}

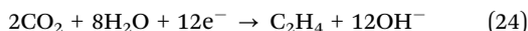


Ji *et al.*⁷¹ reported low cell voltage with a Cu on polymer catalyst in an AEM, while Duong *et al.*¹⁶⁵ achieved high faradaic efficiency with a dendritic Cu catalyst in an AEM. The current densities reported in both studies were similar, so the former was chosen as reference since the reported cell voltage (2.01 V) was about half.

A.4 Direct route

Direct Fischer–Tropsch synthesis (D-FT). Similar to FT synthesis of low olefins from syngas (S-FT), the development of a bifunctional catalyst for direct FT synthesis from CO₂ (reactions (6) and (21)) remains at the research stage (TRL 4).^{168–170} Direct conversion of CO₂ and H₂ to ethylene through FT was also reported using a Fe-K/γ-Al₂O₃ catalyst under 10 bar and 300 °C.¹⁷¹ Do and Kim¹⁷² recently designed a heat-integrated process using this Fe-K/γ-Al₂O₃ catalyst, which was selected as reference. Nevertheless the mass balances provided by Do and Kim¹⁷² are incomplete in that they do not allow quantifying the purge and wastewater streams from D-FT.

CO₂ electroreduction to ethylene (D-eC₂H₄). Similar to CO electroreduction to ethylene, research on direct electroreduction of CO₂ to ethylene (half-cell reactions (24) and (25)) has converged on Cu as the catalyst of choice, with operation under flowing alkaline conditions (TRL 4).^{110,173}



Dinh *et al.*⁷² and Li *et al.*¹¹⁴ both reported ethylene production with high faradaic efficiency and current density under low cell voltage. Dinh *et al.*⁷² devised a multilayer graphite/carbon nanoparticles/Cu/PTFE electrode, while Li *et al.*¹¹⁴ used a Cu-ligand catalyst on an AEM. Since the current densities (1100 A m⁻² and 1390 A m⁻², respectively) are close, the cell voltages (1.78 V and 2.03 V, respectively) became the decisive criterion, making Dinh *et al.*⁷² the chosen reference. Though it is important to note that the current density in this reference was collected in a flow cell with a 7 M KOH electrolyte, so reducing the electrolyte molarity for scaling-up production could incur much greater ohmic losses and increase the cell voltage in turn.

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