



Cite this: *Sustainable Energy Fuels*,
2022, 6, 4749

Evaluation of the potential use of e-fuels in the European aviation sector: a comprehensive economic and environmental assessment including externalities†

Diego Freire Ordóñez,^a Thorsteinn Halfdanarson,^c Caroline Ganzer,^{ac}
Nilay Shah,^a Niall Mac Dowell^{ac} and Gonzalo Guillén-Gosálbez^{id}*^d

The decarbonisation of the transportation sector is key to meeting the climate goals. Whilst the electrification of road passenger transportation is proving to be a viable low-carbon solution in many contexts, a viable pathway towards a decarbonised aviation sector remains opaque. In this context, so-called e-fuels produced via the combination of H₂O, CO₂ and renewable energy may have promise owing to their compatibility with existing infrastructure. Most studies on e-fuels focus only on the economic dimension, neglecting their environmental performance and associated costs. Here, we present a techno-economic evaluation and cradle-to-grave life cycle assessment of Fischer–Tropsch (FT) e-jet fuels produced at different locations in Europe from a range of CO₂ and green H₂ sources to comprehensively assess their potential in aviation, explicitly accounting for externalities. Our results show that e-jet fuel is at present much more expensive (at least 5.4-fold) than its fossil analogue, even when externalities are included (*i.e.*, at least 2.3 fold the current cost of fossil jet fuel). Furthermore, e-jet fuels could exacerbate the damage to human health and ecosystems despite showing lower carbon footprint and resource scarcity impacts than their fossil counterparts. Overall, e-jet fuel could become more economically and environmentally attractive by reducing the cost and impact of CO₂ and green H₂ and, more specifically, the electricity used in their production processes. In this regard, the production plant's location emerges as a critical factor due to the costs associated with balancing the intermittency of site-specific renewables.

Received 31st May 2022
Accepted 27th August 2022

DOI: 10.1039/d2se00757f

rsc.li/sustainable-energy

1. Introduction

Currently, energy consumption increases *ca.* 1% to 2% per year on average,¹ with transport accounting for around 25% of the world's energy demand and 60% of total oil consumption annually.² Thus, the transport sector is responsible for about 20% of the world's carbon emissions.³ Hence, any credible effort to mitigate climate change must also account for emissions from transport. Consequently, a range of different options have been explored, *e.g.*, electrification, H₂, biofuels and synthetic fuels, where the latter two have received particular attention because they could be blended with conventional fuels without changing existing infrastructure. e-Fuels are liquid or gaseous fuels produced from electrolytic H₂ via renewable electricity and captured carbon or nitrogen.⁴ Because

both the power and the feedstocks come from renewable sources,⁵ the so-called renewable electrofuels have the potential to be carbon neutral in terms of greenhouse gas emissions.⁶ However, their energy-intensive nature, high production costs and the need for green H₂ and concentrated CO₂ streams constitute the main barriers to widespread deployment.^{7–10}

According to Hänggi *et al.*,¹¹ when it comes to passenger automobiles, electric batteries are deemed more sustainable and cleaner than fossil fuels. However, long-distance transportation by land and sea and commercial air travel would require unique batteries unable to currently match present requirements, such as a high gravimetric energy density, a lightweight, a high degree of autonomy, or a short refuelling time.^{12,13} As a result, electrofuels may find use in transport applications where batteries are not suitable.^{6,14}

According to Åkerman *et al.*,⁷ there are three potential aviation fuel substitutes for conventional fossil fuels: drop-in biofuels, H₂, and drop-in electrofuels. The supply of sustainable biomass is inherently limited and owing to the increasingly pressing need to remove CO₂ from the atmosphere, biomass resources will likely be employed in this context. Whilst H₂ could be produced at a lower cost than electrofuels, it is not

^aThe Sargent Centre for Process Systems Engineering, Imperial College London, UK

^bInstitute for Applied Sustainability Research, Quito, Ecuador

^cCentre for Environmental Policy, Imperial College London, UK

^dInstitute for Chemical and Bioengineering, ETH Zürich, Switzerland. E-mail: gonzalo.guillen.gosalbez@chem.ethz.ch

† Electronic supplementary information (ESI) available. See <https://doi.org/10.1039/d2se00757f>



Table 1 Literature review of the production of liquid carbon-based electrofuels for the aviation sector. Most studies focus only on GW indicators and do not consider an explicit valuation of environmental impacts

Source	Main focus	Economic assessment		Environmental assessment			Economic valuation of environmental impacts (externalities)		Uncertainty analysis	
		Yes	No	Only GW indicators	GW + other indicators	No	Yes	No	Yes	No
Åkerman <i>et al.</i> ⁷	Analysis of different scenarios and alternatives towards a more sustainable long-distance travel in Sweden in line with the Paris agreement	×				×		×	×	
Alhyari <i>et al.</i> ¹⁵	Assessment of the environmental impact of power-to-liquid (PtL) processes by applying LCA		×		×			×		
Goldmann <i>et al.</i> ⁶	Technical evaluation of <i>n</i> -octane, methanol, methane, H ₂ and ammonia as possible substitutes for conventional aviation fuels		×			×		×		×
Kieckhäfer <i>et al.</i> ¹⁶	Application of a system dynamics approach to assessing the possibility for alternative jet fuel uptake in aviation and the extent to which alternative fuels can help meet CO ₂ reduction targets	×		×				×	×	
Schemme <i>et al.</i> ¹⁷	Techno-economic analysis of power-to-fuel (PtF) processes based on the same harmonised premises and boundary conditions	×				×		×	×	
Schmidt <i>et al.</i> ¹⁸	Review of the environmental and techno-economic performance of state-of-art PtL production processes for aviation	×			×			×	×	
Schmidt <i>et al.</i> ¹⁹	Review of technical, economic, and environmental aspects and potential of PtL processes for renewable jet fuel production	×			×			×	×	
Sherwin <i>et al.</i> ²⁰	Economic viability analysis of electrofuels synthesised from CO ₂ captured using direct air capture (DAC), and H ₂ produced <i>via</i> water electrolysis, predominantly powered by solar or wind energy	×		×				×	×	
Trinh <i>et al.</i> ²¹	Development of an optimisation-based model to assess the impact of Sweden's carbon pricing, blending mandates, and penalty charge on renewable jet fuel production	×		×				×	×	

a drop-in option for the existing fleet, which may be expected to remain in service for some time. Therefore, this study focuses on electrofuels with the potential to be used as aviation fuels.

Recent literature studied the potential benefits and feasibility of electrofuels in the aviation sector, as shown in Table 1. Most of these studies focus only on the techno-economic dimension. Only a few assess their environmental performance, but often in terms only of global warming (GW) impacts. However, as discussed elsewhere,^{8–10} a full assessment of the potential of electrofuels to contribute to a net-zero emission society should consider both their techno-economic viability and environmental performance at an equivalent level of granularity. Moreover, drawing a comprehensive picture of environmental performance requires accounting for impacts beyond climate change. The latter can be achieved *via* a comprehensive life-cycle assessment (LCA). Specifically, electrofuels are currently more expensive but could be less polluting, depending on the carbon intensity of the energy used in their production, than their fossil analogues.²² However, they may impose significant indirect impacts on human health and the environment. Consequently, a fair comparison requires accounting for other impacts beyond climate change, which could be done by computing the monetised value of the life cycle environmental impacts.

Here, we present a comprehensive techno-economic and environmental study of Fischer-Tropsch (FT)-electrofuels with the potential to replace fossil jet fuel, covering a wide range of LCA metrics, externalities and several locations.

We focus on e-jet fuel obtained from CO₂ captured from different sources and green H₂ from solar, wind, and hybrid solar and wind-powered water-electrolysis systems. Our study considers various locations in Europe, each with different availability of solar and wind resources, which affects the cost of electricity and H₂ storage, and thus the final cost of H₂ and e-jet fuel.^{23,24} The plant integrates solar PV and wind systems, a proton exchange membrane electrolyser (PEMEL) and its corresponding balance of plant (BOP), alongside batteries and H₂ storage tanks (or salt caverns). It also includes a reverse water-gas shift (rWGS) reactor, an FT reactor, a hydrocracking (HC) reactor, a burner, and a train of flash separators. The LCA undertaken here is based on the ISO 14040 series (2006) and the ReCiPe 2016 damage model, covering 22 midpoint and three endpoint indicators that are monetised according to specific economic penalties.²⁵

We find that, at present, e-jet fuel is more expensive than fossil jet fuel by a factor of 5.4–15.1, even when externalities are accounted for. For e-jet fuels to be cost-competitive with conventional jet fuel, a carbon tax of 1125 USD/t_{CO₂-eq} would need to be imposed, which is deemed unrealistic. Moreover, e-jet fuel leads to burden-shifting as it worsens human health and ecosystem quality compared to fossil jet fuel. Overall, our work highlights the need to account for impacts beyond climate change in assessing electrofuels to minimise the potential collateral damage of low carbon transition pathways.

The paper is organised as follows: the methodology encompassing the model description, the LCA and techno-economic assessment, including the monetisation of endpoint impacts and the risk analysis, is outlined first. Following that, the results

are presented, comparing e-jet fuel with conventional jet fuel. Finally, we summarise the primary findings and provide recommendations for further research.

2. Methodology

We consider five plant locations, namely the United Kingdom (GB), Germany (DE), France (FR), Spain (ES) and Italy (IT), given their high jet fuel demand in the European region (see Fig. 1).²⁶ A summary of the scenarios assessed for each country, highlighting the H₂ and CO₂ sources and H₂ storage media, is shown in Table 2.

2.1. Model description

The production process, which corresponds to the foreground system exchanging mass and energy with the techno- and eco-spheres, was simulated in Aspen Plus v11 (ref. 27) based on the literature.²⁸ The model includes the pure components H₂, CO₂, H₂O, CO, and all the n-alkanes from C₁ to C₃₀, which were selected from the APV110.PURE37 databank. Due to its suitability for hydrocarbons,²⁹ the Peng–Robinson equation of state with Boston–Mathias alpha function (PR-BM) was chosen to estimate thermodynamic properties.

Fig. 2 provides the process flowsheet. CO₂ and H₂ react in an rWGS reactor to produce syngas, which is sent to a low-temperature FT reactor. The FT gases are then separated and upgraded *via* a flash separation train, whereas FT waxes undergo HC in a hydrocracker. The resulting products enter a final flash separator that purifies the e-jet fuel. In the next sections, we describe each step of the production process in detail.

2.1.1. Feedstocks: green H₂ and captured CO₂

Green H₂. Green H₂ is produced from water electrolysis using renewable electricity, according to eqn (1).

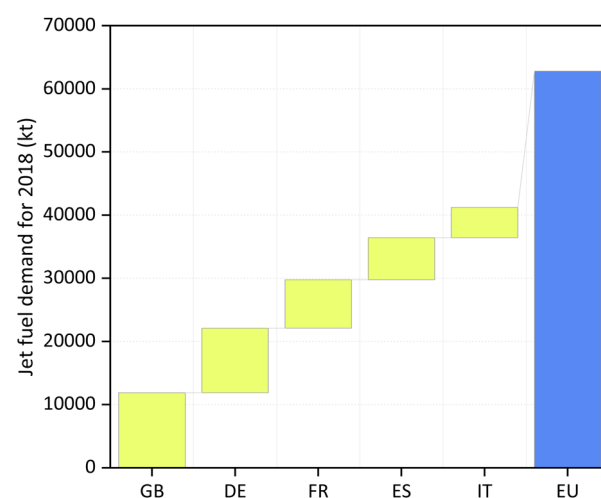


Fig. 1 European demand for jet fuel in 2018. The European Union's (EU) jet fuel demand for 2018 was 62 800 kt. The sum of the demands of the United Kingdom (GB), Germany (DE), France (FR), Spain (ES) and Italy (IT), for the same year, was equivalent to approximately 65% of the EU's total demand.²⁶

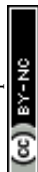
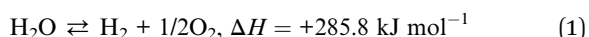


Table 2 H₂ and CO₂ sources and H₂ storage media for each e-jet fuel production scenario

Scenario	H ₂ source	H ₂ storage media	CO ₂ source
1	Hybrid solar and wind-based electrolysis	Type I tanks	Coal-based power plant (C-PP)
2	Hybrid solar and wind-based electrolysis	Salt caverns	Coal-based power plant (C-PP)
3	Hybrid solar and wind-based electrolysis	Type I tanks	Natural gas-based power plant (NG-PP)
4	Hybrid solar and wind-based electrolysis	Salt caverns	Natural gas-based power plant (NG-PP)
5	Hybrid solar and wind-based electrolysis	Type I tanks	Direct air capture based on a high temperature, liquid solvent system (DAC-HT)
6	Hybrid solar and wind-based electrolysis	Salt caverns	Direct air capture based on a high temperature, liquid solvent system (DAC-HT)
7	Hybrid solar and wind-based electrolysis	Type I tanks	Direct air capture based on a low temperature, solid sorbent system (DAC-LT)
8	Hybrid solar and wind-based electrolysis	Salt caverns	Direct air capture based on a low temperature, solid sorbent system (DAC-LT)



Other works assumed a fixed location to compute the H₂ cost. In contrast, here, we applied an existing modelling framework, *i.e.*, the gAWE model,²⁴ to estimate the electrolytic H₂ cost in various locations, thus explicitly considering the location-specific availability of renewable energy. The original model, which focused on green H₂ from solar power, was extended to account for wind as an alternative energy source. The H₂ production plant encompasses solar PV and wind

facilities and the PEMEL system. The latter includes its BOP, and electricity and H₂ storage systems, *i.e.*, batteries and Type I tanks (or salt caverns), respectively. We consider the PEMEL technology given its high TRL and market availability.³⁰ The fluctuating nature of renewables is taken into account by estimating the cost of storing electricity and H₂ to guarantee a continuous supply of H₂ to the fuel production plant, which substantially contributes to the NPC.^{23,24}

The model, which is formulated as a mixed-integer linear programming (MILP) model, evaluates the cost-based optimal

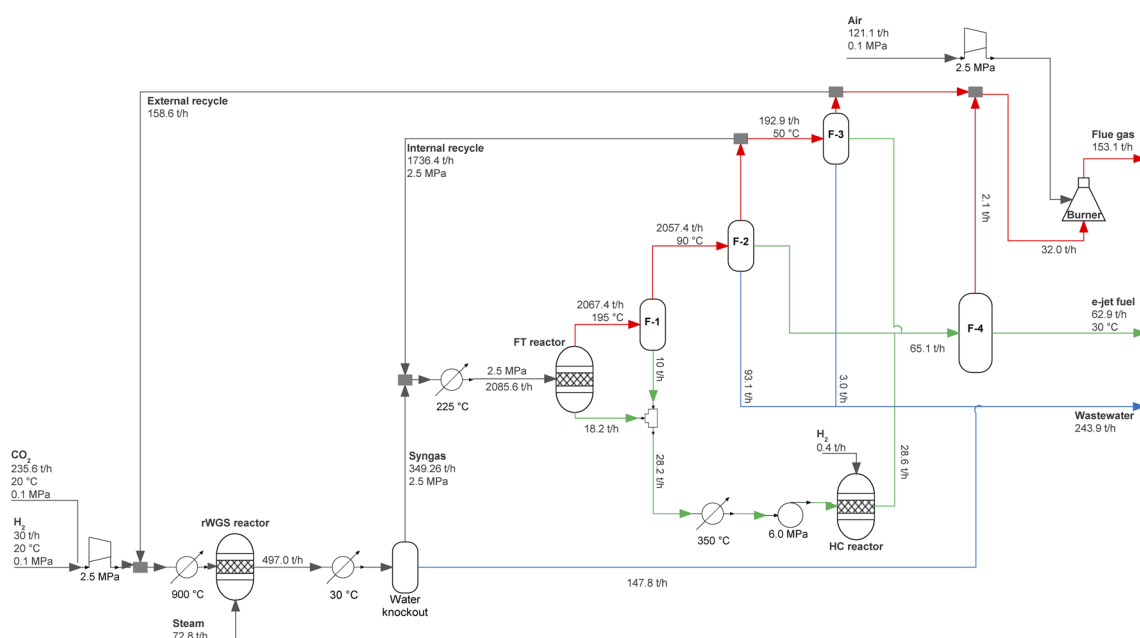


Fig. 2 Flowsheet of the production process of e-jet fuel. Syncrude is produced in a low-temperature FT reactor from the syngas obtained from CO₂ and H₂. e-Jet fuel is then obtained from the separation of syncrude, according to the operating conditions depicted in the figure. A hydrocracker is implemented to process FT waxes and increase the fuel yield. The colour coding for the streams after the FT reactor is as follows: gaseous streams (red), liquid streams (orange), and wastewater (blue).



H₂ production configuration considering raw materials and energy consumption, equipment sizing, heat integration requirements, and electricity and H₂ storage capacities for a given hourly-based production target. We consider the production of 1 GW_{LHV} of H₂ (ref. 31–33) as input for the chemical plant plus an additional amount for the hydrocracker (see Fig. 2). We evaluated the production of green H₂ in the five European countries for six different configurations; namely, (i) solar-powered PEMEL with Type I tanks for H₂ storage, (ii) solar-powered PEMEL with salt caverns for H₂ storage, (iii) wind-powered PEMEL with Type I tanks for H₂ storage, (iv) wind-powered PEMEL with salt caverns for H₂ storage (iii) hybrid solar and wind-powered PEMEL with Type I tanks for H₂ storage, and (v) hybrid solar and wind-powered PEMEL with salt caverns for H₂ storage. The MILP model was solved using GAMS 32.1.0 and the solver CPLEX 12.10.0.0. The hourly-based solar PV and wind capacity factors (CF) were taken from renewables.ninja^{34,35} for 2018. The model input data can be seen in Appendix C, Table C-2,† while additional details can be found in ref. 24.

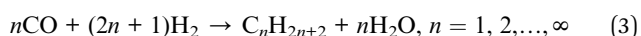
Captured CO₂. The concentrated CO₂ required for electrofuels production³⁶ entails a different environmental impact and can be obtained at various prices depending on the source. Thus, we evaluated three potential sources of CO₂, namely CO₂ captured from coal power plants (C-PP), natural gas power plants (NG-PP) and CO₂ recovered directly from the atmosphere – direct air capture (DAC). The DAC technology is characterised by either a high temperature, liquid solvent system (DAC-HT) or a low temperature, solid sorbent system (DAC-LT). For CO₂ from power plants, a system expansion strategy is employed to deal with the multi-functionality of electricity production. Data for the capture processes were taken from existing literature (Appendix B, Table B-4†).

2.1.2. Syngas production from CO₂ and H₂: rWGS reaction. Syngas is a gaseous mixture primarily composed of H₂, CO, and traces of CO₂. It is produced by the rWGS reaction taking place at a high temperature:³⁷



In our case, we assume that syngas is obtained from CO₂ and H₂ at 900 °C using a nickel-based catalyst.^{38,39} This catalyst was selected due to its high activity and low cost compared to noble metal-active phase catalysts.⁴⁰

2.1.3. Syncrude production from syngas: Fischer–Tropsch (FT) process. The FT process consists of polymerisation reactions whereby CO₂ and H₂ are converted into liquid hydrocarbons.⁴¹ For the model, we consider the hydrogenation of CO to form *n*-paraffins as the dominant reaction:



We assumed here that the product distribution of the FT reactions follows the Anderson–Schulz–Flory (ASF) distribution:⁴¹

$$w_n = n(1 - \alpha)^2 \alpha^{n-1} \quad (4)$$

where w_n denotes the weight fraction of hydrocarbons with carbon number (or chain length) n , and α the chain growth probability.

Iron and cobalt are the preferred catalysts for FT industrial applications.⁴² For this study, we consider a low-temperature FT process (180–250 °C)⁴³ using a cobalt catalyst^{44,45} ($\alpha = 0.90$ – 0.95).⁴⁶ This is because, for low-temperature FT processes implementing cobalt-based catalysts, only unbranched alkanes (C_{*n*}H_{2*n*+2}) are often obtained as products.¹⁷ As far as methane is concerned, it is well known that its selectivity is higher than predicted by the ASF distribution.⁴¹ To take this into account, we followed the approach by König *et al.*,^{39,47} which considers a correction factor for methane (methane yield correction, w_{CH_4} , of 0.04 for $\alpha = 0.95$) based on an experimentally obtained FT product distribution.⁴⁸ Additional FT design parameters can be found in Appendix A, Fig. A-1.†

2.1.4. FT wax hydrocracking (HC). HC is a catalytic cracking process whereby the C–C bonds in aromatics and olefins, *i.e.*, long hydrocarbon chains, are broken by the addition of H₂ to yield naphthenes and alkanes.^{49,50}

Hydrocracking of FT waxes, *i.e.*, long paraffinic chains (C₂₁),⁵¹ provides high-end middle distillates with low sulphur and aromatics content.⁵² Regarding the catalysts, NiMo and NiW supported on solid acids are particularly well suited for producing middle distillates.⁵² However, Pd and Pt-loaded catalysts, which have high hydrogenation/dehydrogenation activity for heavy hydrocarbon cracking, could also be used owing to the sulphur-free nature of FT waxes.^{51,53,54} Thus, we considered a platinum-based catalyst supported on amorphous silica-alumina in this work.^{53,55} The HC product distribution is based on Calemme *et al.*⁵⁴ (see Appendix A, Fig. A-1†).

2.1.5. Product separation and upgrading. In order to refine the properties of the e-jet fuels, FT gases and HC products are fed to a train of flash separators whose operating conditions are adjusted to yield fuels with properties as close as possible to those of commercial jet fuels. Hence, four flash separators were implemented, as shown in Fig. 2.

2.1.6. Heat integration (HI). Process integration focused on minimising mass and energy consumption can be applied to improve the economic and environmental performance.⁵⁶ In this study, once the flowsheet is built and utilities are assigned in Aspen Plus, pinch analysis was conducted *via* Aspen Energy Analyzer v11 (ref. 57) to determine the potential energy savings of HI. For this purpose, the software generates the composite curve and evaluates the energy targets. In addition, a heat recovery approach temperature (HRAT) of 10 °C was considered.

2.2. Life-cycle assessment (LCA)

Here, we applied LCA according to the guidelines given by Bruijn *et al.*⁵⁸ to quantify environmental impacts. These are linked to life cycle activities connected to the production and utilisation of fossil jet fuel and e-jet fuel, including, in the latter case, green H₂ production. The LCA models were implemented in SimaPro v9.0 (ref. 59) interfacing with Ecoinvent 3.5. The modelling approach and assumptions are explained below.

2.2.1. Goal and scope definition. We considered a cradle-to-grave approach, including both the life cycle emissions



from the production and combustion of the fuels. The functional unit is 1 kg of e-jet fuel produced and combusted. We chose a mass-based functional unit following recommended guidelines for carbon capture and utilisation (CCU).⁶⁰ We disregarded the environmental impacts associated with the manufacturing of the PEMEL since their contribution to the total impacts was found to be low.^{61–63}

2.2.2. Life-cycle inventory (LCI). The LCI combines data from the foreground system, *i.e.*, mass and energy flows in the Aspen simulation, and background system, *i.e.*, surrounding processes providing inputs to the main process. Information on the foreground system was gathered from the gAWE model runs (for the H₂ feedstock) and the model simulation in Aspen Plus. In contrast, data from the background system was retrieved from Ecoinvent v3.5 (see Appendix B†). The LCI for fossil jet fuel (BAU) production was collected directly from Ecoinvent v3.5. The emission factors to account for fuels utilisation were taken from Nojoudi *et al.*,⁶⁴ as shown in Appendix B, Table B-2.† Allocation at the point of substitution (APOS) was applied because it includes both the impacts of the production and treatment processes.⁶⁵

2.2.3. Environmental impact assessment (EIA). On the basis of the hierarchist cultural perspective,⁶⁶ the ReCiPe 2016 framework⁶⁷ was applied as implemented in SimaPro. We focused our analysis on global warming potential (GWP) and the three endpoint indicators, *i.e.*, human health (HH), ecosystem quality (EQ), and resource scarcity (RS), obtained by aggregating different midpoint indicators, as follows:

- HH (DALY): global warming (HH), stratospheric ozone depletion, ionising radiation, ozone formation (HH), fine particulate matter formation, human carcinogenic toxicity, human non-carcinogenic toxicity, and water consumption (HH).
- EQ (species yr): global warming (terrestrial ecosystems), global warming (freshwater ecosystems), ozone formation (terrestrial ecosystems), terrestrial acidification, freshwater eutrophication, marine eutrophication, terrestrial ecotoxicity, freshwater ecotoxicity, marine ecotoxicity, land use, water consumption (terrestrial ecosystems), and water consumption (aquatic ecosystems).
- RS (USD₂₀₁₃): mineral resource scarcity and fossil resource scarcity.

2.3. Economic analysis

2.3.1. Net production cost (NPC)

Green H₂. The NPC of green H₂ was calculated from the capital expenditures (CAPEX) of the solar PV, wind and PEMEL systems and storage options, based on the design capacities obtained from the gAWE model. The OPEX was considered negligible since raw materials (H₂O, sunlight and wind) are freely available, utilities (electricity) are modelled explicitly and fixed OPEX expenditures, such as labour cost, are low as well, relative to other costs.²⁴ The CAPEX corresponds to the fixed capital investment (FCI) of the H₂ production plant. The working capital (WC) was assumed to be 10% of the total capital

investment (TCI).²⁹ The TCI, *i.e.*, the sum of the FCI and WC, was then annualised according to the equation below:

$$ACC = FCI \left(\frac{i(1+i)^t}{(1+i)^t - 1} + \frac{i}{9} \right) \quad (5)$$

where ACC represents the annualised capital cost, *i*, the interest rate and *t*, the plant economic life. The second term of the equation represents the WC.

Finally, the NPC of green H₂ was calculated by dividing the ACC by the annual production (AP), as shown below:

$$NPC = \frac{ACC}{AP} \quad (6)$$

The NPC of H₂ is expressed in USD per kg.

The cost input data required by the model can be seen in Appendix C, Table C-2.†

e-Jet fuels. The NPC of the e-jets was obtained from the CAPEX and OPEX associated with the fuel production plant. These costs were calculated using the mass and energy flows obtained from the model simulation in Aspen Plus. The purchased cost (PC) of the conventional equipment units, *i.e.*, heat exchangers, compressors, separators, and pumps, was calculated using the correlations given by Sinnott and Towler.⁶⁸ On the other hand, the PC of special equipment units, *i.e.*, the rWGS reactor, FT reactor, hydrocracker and burner, was obtained from eqn (7) using the parameters listed in Appendix C, Table C-3.† All costs, including raw materials, utilities, products, and equipment, were updated to 2018 using price indices, as shown below. The chemical engineering plant cost index (CEPCI)⁶⁹ was applied to all equipment units. In contrast, the World Bank Commodity Price Data (The Pink Sheet)⁷⁰ was employed in raw materials and utilities.

$$PC = PC_{\text{ref}} \left(\frac{S}{S_{\text{ref}}} \right)^D \times \left(\frac{CEPCI_{2018}}{CEPCI_{\text{ref}}} \right) \quad (7)$$

In the equation above, PC represents the purchased cost of the equipment, PC_{ref} denotes the same purchased cost expressed in monetary units of the reference year, *S* indicates the actual size of the equipment, *S*_{ref} reflects the reference capacity of the equipment, and *D* is a scaling factor. The CEPCI₂₀₁₈ is assumed to be equal to 603.1.⁷¹ According to the Pink Sheet, for the same year, the energy index is 85.48 and the non-energy index 83.72.⁷⁰ The former is used to update utility costs and the latter to update raw material costs.

The FCI of the fuel production plant and its total operating cost (TOC) were estimated following the guidelines given by Albrecht *et al.*²⁹ based on the data presented in Appendix C, Tables C-3 and C-4.† The FCI includes the purchased equipment, installation, and further capital requirements in the construction phase. The TOC encompasses raw materials and utilities, costs/revenues from by-products, labour costs, maintenance costs, and other indirect costs.²⁹ The ACC was obtained from eqn (5), with the data in Appendix C, Table C-1.† Lastly, the total annualised cost (TAC) and the NPC of e-jet fuel were calculated as follows:



$$\text{TAC} = \text{TOC} + \text{ACC} \quad (8)$$

$$\text{NPC} = \frac{\text{TAC}}{\text{AP}} \quad (9)$$

The NPC of e-jet fuel expressed in USD per $L_{\text{fossil jet fuel-equivalent}}$ was converted to USD per t per km, *i.e.*, fuel cost per load (in tonnes) and distance flown (in kilometres). We considered a conversion factor of 0.325 L per t per km for 2018.^{72–74}

2.3.2. Abatement cost of carbon emissions. For e-jet fuel, the cost of carbon avoided, *i.e.*, the carbon tax required to bring e-jet fuel up to the cost level of their fossil-based counterpart, was calculated from eqn (10):

$$\text{cost of carbon avoided} = \left| \frac{\text{NPC}_{\text{e-jet fuel}} - \text{NPC}_{\text{fossil jet fuel}}}{\text{GWP}_{\text{e-jet fuel}} - \text{GWP}_{\text{fossil jet fuel}}} \right| \quad (10)$$

here, GWP and NPC represent the global warming potential and net production cost, respectively. The GWP of the fuels ($\text{kg}_{\text{CO}_2\text{-eq}} \text{kg}_{\text{fuel}}^{-1}$) is obtained from the results of the LCA at the midpoint level.

2.3.3. Monetisation of LCA endpoints (externalities). The monetisation of the LCA impacts allows the economic quantification of environmental damages. As done elsewhere,^{8,75–77} the environmental impact indicators at the endpoint level were monetised according to the following penalties,²⁵ which values were updated to 2018 using price indices.

- HH. A value of 7.4×10^4 EUR₂₀₀₃ per 1 DALY is assumed to be the willingness-to-pay value for a life year with a minor risk increase from involuntary exposure.

- EQ. A value of 9.5×10^6 EUR₂₀₀₃ per 1 lost species/year is derived based on the share of our well-being that we are ready to give up to protect ecosystems.

- RS. A value of 8.62×10^{-1} EUR₂₀₀₃ per 1 USD₂₀₀₀ is obtained by converting USD₂₀₀₀ to EUR₂₀₀₃ considering inflation and the International Monetary Fund mid-year SDR exchange rate for 2003.

2.3.4. Uncertainty analysis. Monte Carlo simulations were conducted to study the impact on the NPC of key uncertainties. To this end, 10 000 independent simulations were conducted for the cheapest alternative of each scenario in Table 2, varying the H₂, CO₂ and grid electricity costs. For each simulation, a normal distribution with a standard deviation of 20% with respect to the current costs was assumed, as suggested elsewhere.⁶⁸

3. Results and discussion

In this section, the results of the process models are discussed first, to then focus on the environmental and economic results.

3.1. Process models

A summary of the design capacities of the equipment in the H₂ production plant is given in Table 3, whereas the mass and energy flows are shown in Table 4.

Focusing first on green H₂, we find that in most of the locations, *i.e.*, GB, FR, ES and IT, the design capacity of PV exceeds that of wind systems. This is because, in these locations, the annual CF for wind is higher than the one for solar PV systems.

Regarding PEMELs, the lowest design capacity corresponds to wind-solar hybrid systems due to the combined use of solar and wind resources. The model chooses H₂ tanks as storage option for all locations due to the current prohibitively high cost of Li-ion batteries, *ca.*, 420 USD/kWh.⁷⁸ The required capacity of

Table 4 Summary of mass flows and heat integration results for a 1 GW_{LHV} H₂ input plant

Mass flows		
Streams	Value	Unit
Inputs		
CO ₂	235.6	t h ^{−1}
H ₂	30.4	t h ^{−1}
H ₂ O	72.8	t h ^{−1}
Air	121.1	t h ^{−1}
Outputs		
e-Jet fuel	62.91	t h ^{−1}
Flue gas	153.102	t h ^{−1}
Wastewater	243.93	t h ^{−1}
Energy flows		
Streams	Value	Unit
Inputs		
Electricity	122.2	MW
Cooling water	246.1	MW
Outputs		
e-Jet fuel	764.2	MW

Table 3 Design capacities of H₂ production plant equipment required for continuous production of green H₂. A total production of 30.4 t_{H₂} h^{−1} is assumed for solar-based H₂, wind-based H₂, and hybrid solar and wind-based H₂ at each location

Equipment	GB			DE			FR			ES			IT		
	Solar H ₂	Wind H ₂	Hybrid H ₂	Solar H ₂	Wind H ₂	Hybrid H ₂	Solar H ₂	Wind H ₂	Hybrid H ₂	Solar H ₂	Wind H ₂	Hybrid H ₂	Solar H ₂	Wind H ₂	Hybrid H ₂
PV system (GW)	23.5	—	3.3	20.7	—	6.0	20.2	—	4.8	15.1	—	3.1	15.7	—	6.5
Wind system (GW)	—	5.3	3.2	—	21.6	7.4	—	9.8	4.2	—	10.2	6.1	—	9.2	2.7
PEMEL (t _{H₂} h ^{−1})	159.2	56.9	49.1	173.0	76.1	69.4	161.5	59.7	58.5	148.7	68.0	52.2	147.4	84.4	69.5
H ₂ storage tanks or caverns (t)	54 077	5342	3227	17 823	8086	6653	33 870	4670	3592	4507	5675	5978	6236	10 048	5196
Electricity storage (GWh)	—	—	—	—	—	—	—	—	—	—	—	—	—	—	—



the storage tanks is adjusted by the model depending on the size of the solar PV, wind, and electrolysis systems.

Some of the most important properties of fossil jet fuel, and the corresponding properties estimated from the simulation for e-jet fuel, are shown in Appendix A, Table A-1.†

The energy analysis of the production process shows that the heating demand, *i.e.*, 363.5 MW, can be entirely fulfilled by heat integration. Likewise, the cooling demand can be reduced by

60% compared with the non-integrated process (*i.e.*, from 609.7 MW to 246.1 MW). The electricity required for the fuel production plant, *i.e.*, 122.2 MW, is assumed to be taken directly from the grid to avoid expensive electric batteries. As grid electricity comes from different renewable and non-renewable sources, there are different amounts of carbon embedded in this electricity, depending on the location of the plant.

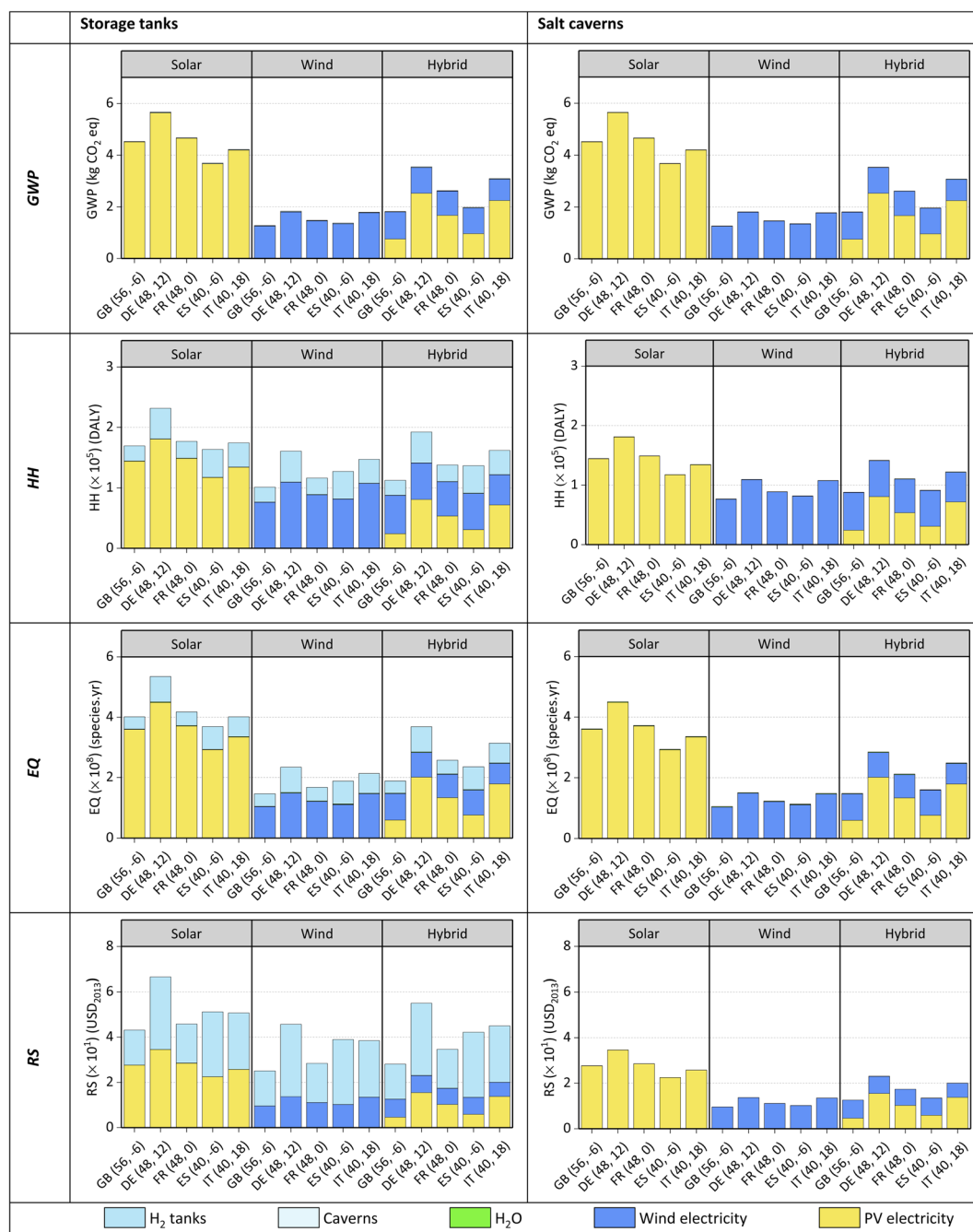


Fig. 3 Green H₂: LCA at the endpoint level + G: LCA at the endpoint level + WP. GWP: global warming potential, HH: human health, EQ: ecosystem quality, RS: resource scarcity. The figure shows the environmental impacts of the production of green H₂ from solar, wind and solar-wind (hybrid) power in the United Kingdom (GB), Germany (DE), France (FR), Spain (ES) and Italy (IT), according to the ReCiPe 2016 LCA methodology. Most of the environmental impact of H₂ comes from the electricity used for the electrolysis of water. For the scenarios assessed, wind H₂ and solar H₂ show the lowest and the highest impacts, respectively. Furthermore, tanks show higher impacts than salt caverns.



3.2. Life-cycle assessment (LCA)

3.2.1. Analysis at the endpoint level + GWP: green H₂. The LCA results are shown in Fig. 3.

We start by analysing the environmental footprint of green H₂, the main contributor to the impact embodied in the e-fuel. Most of the environmental impacts associated with green H₂ come from the electricity used in water electrolysis. The reason behind this is

the fossil energy embodied in solar panels and wind turbines. However, the carbon footprint of solar PV and wind power is expected to decrease by 2050 compared to today's values by approximately 90% and 70%, respectively.⁷⁹ These reductions would decrease the carbon footprint of electrolytic H₂ substantially.

Overall, H₂ from wind power outperforms both H₂ from solar power and H₂ from hybrid power in GWP and all endpoint categories.

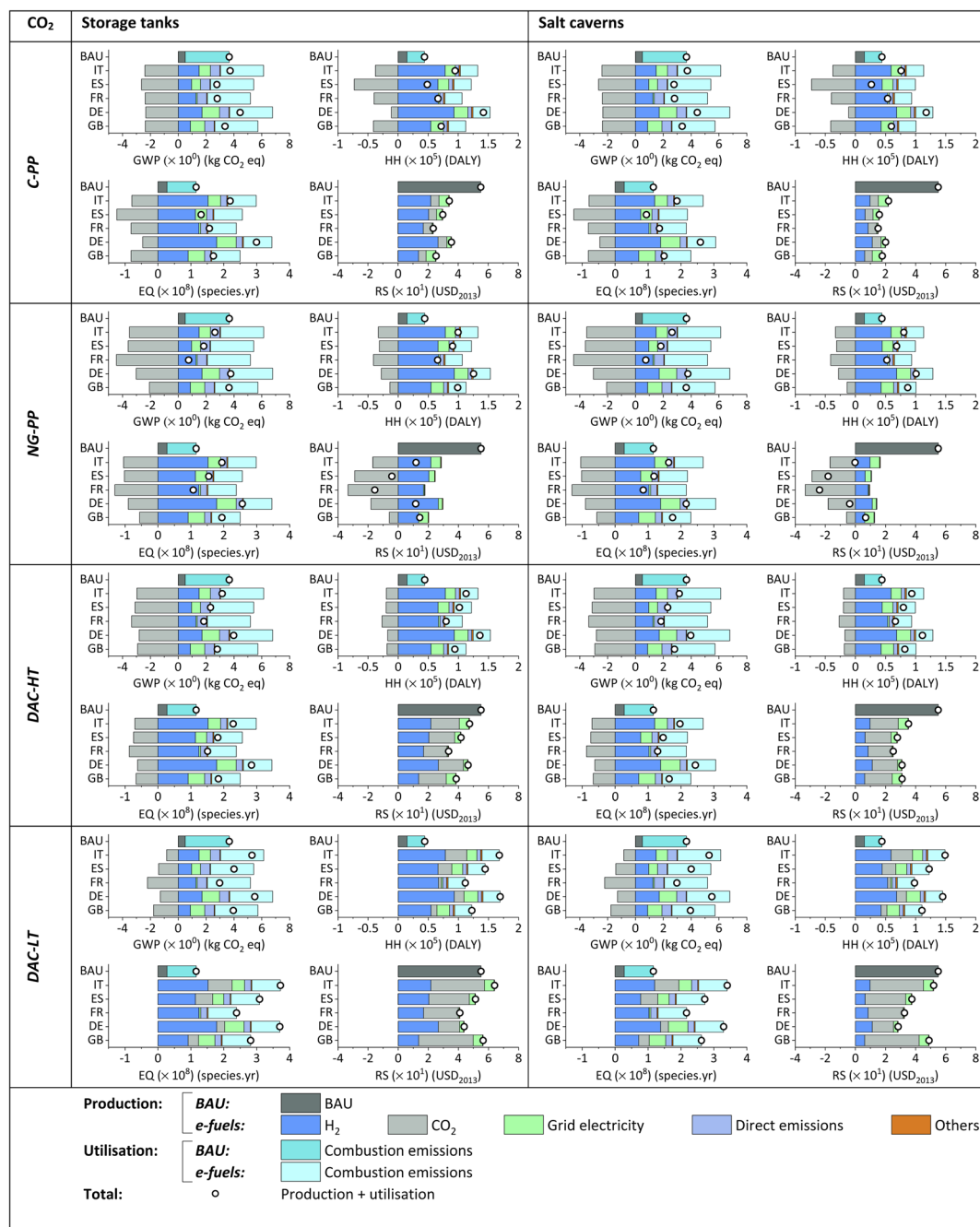
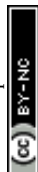


Fig. 4 e-Jet fuel: LCA at the endpoint level + GWP. GWP: global warming potential, HH: human health, EQ: ecosystem quality, RS: resource scarcity. The environmental impacts of the production and utilisation of e-jet fuel from different sources of CO₂ and green H₂ from hybrid power are assessed based on the ReCiPe 2016 LCA methodology, for different locations: United Kingdom (GB), Germany (DE), France (FR), Spain (ES) and Italy (IT). The major contributors to the environmental impact of e-jet fuels are H₂, CO₂ and combustion emissions. There is burden-shifting, as e-jet fuel generally shows better performance in RS compared to fossil jet fuel at the expense of worsening HH and EQ.



Concerning H_2 storage, tanks generate more impacts on HH, EQ and RS than salt caverns for all locations. For GWP, the impacts of Type I tanks and salt caverns are negligible compared to the electricity needed for water electrolysis. The environmental impact of Type I tanks is most notable in the RS endpoint category due to the use of aluminium alloy. For the modelling of salt caverns, only the impacts of compressing and drying H_2 have been considered, which happen to be low in the categories analysed. Finally, although salt caverns would be environmentally better than Type I tanks, their availability could become an issue. Despite its potential for H_2 storage in Europe, subsurface salt structures should meet strict technical criteria related to size, land availability and storage capacity,^{80,81} narrowing down the available options substantially.

3.2.2. Analysis at the endpoint level + GWP: e-jet fuel. Fig. 4 shows the GWP and endpoint-level results of the environmental assessment of the production and utilisation of e-jet fuel at each location, based on the scenarios described in Table 2. The results of the midpoint-level assessment are shown in Appendix D.† As seen, H_2 from solar-wind hybrid systems is selected in all scenarios as it is the cheapest alternative (see Fig. 5).

When comparing e-jet fuels with the BAU alternative, burden-shifting, *i.e.*, some impacts become milder at the expense of worsening others, can be observed at both the endpoint and midpoint levels. Overall, there is burden-shifting toward the HH and EQ impact categories at the endpoint level. However, when using H_2 stored in caverns, e-jet fuel can outperform the BAU in all the impact categories for specific locations, *i.e.*, ES, when CO_2 from C-PP is utilised as feedstock. Regarding GWP, e-jet fuels outperform the BAU in most locations when CO_2 from power plants and DAC-HT are used. Nevertheless, the production of e-jet fuel in DE does not decrease in GWP compared to its fossil analogue, mainly due to the current environmental burden associated with green H_2 . Similarly, e-jet fuel from CO_2 from DAC-LT would only be beneficial in FR.

In terms of the CO_2 source, e-jet fuels from (DAC-HT) CO_2 and (DAC-LT) CO_2 show a higher environmental impact than (NG-PP) CO_2 and (C-PP) CO_2 on HH, EQ and RS. This means that, owing to their energy intensity,^{82,83} the DAC processes operated by both high and low-temperature heat perform worse than point source carbon capture. However, the high-carbon intensity energy currently in use is expected to be progressively decarbonised over the next few years.^{79,84} Moreover, it is worth noting that CO_2 capture through DAC processes, unlike CO_2 capture from power plants, aims to close the carbon loop. Therefore, a large global deployment of DAC technologies is expected in order to achieve the environmental objectives foreseen for the coming decades.

Lastly, from the LCA results of green H_2 , e-jet fuel produced from H_2 stored in salt caverns shows lower environmental impacts in all the categories than when using storage tanks, regardless of the location.

3.3. Economic analysis

3.3.1. Net production cost (NPC)

NPC: green H_2 . The NPC of green H_2 is presented in Fig. 5.

The storage of H_2 in salt caverns is cheaper than in tanks, *ca.* 72%. H_2 from hybrid energy is always the cheapest option, *i.e.*, 32–82% and 8–36% cheaper than solar-based H_2 and wind-based H_2 , respectively. Regarding the cost breakdowns, the major contributors towards the NPC of H_2 are the PV systems and/or wind systems (36–84%) except for solar H_2 in GB, where the most significant investment would be in H_2 tanks (*ca.* 52% of the NPC).

NPC: e-jet fuel. The NPC of e-jet fuel is presented in Fig. 6. H_2 from hybrid solar and wind is considered for all scenarios, as in the LCA. The CAPEX and OPEX breakdowns are shown separately for the cheapest option of each scenario (marked with an *).

H_2 stored in salt caverns is cheaper than H_2 stored in tanks. Consequently, e-jet fuels from the former are thereby cheaper

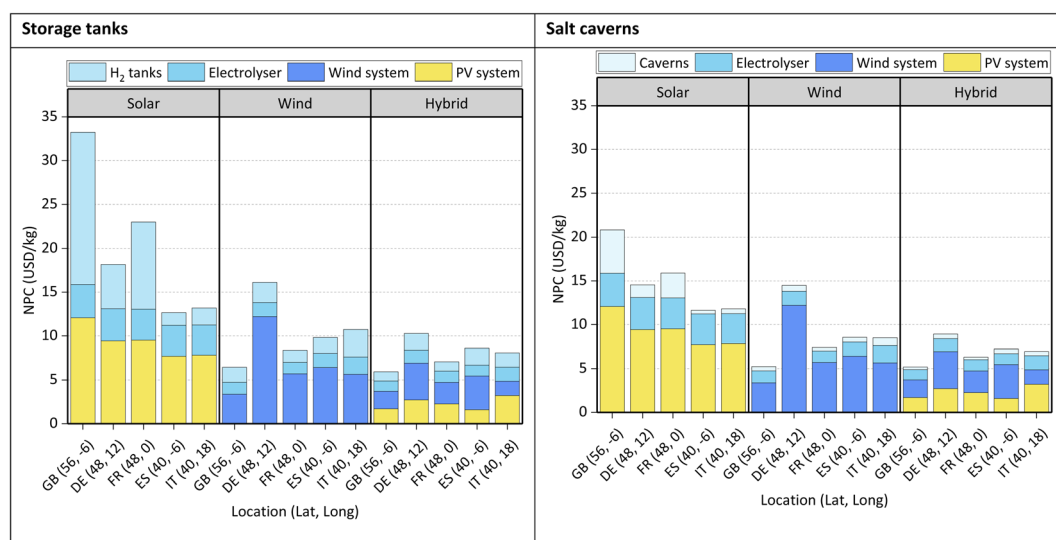


Fig. 5 Green H_2 : NPC breakdown. The NPC of green H_2 from solar energy is higher than that from wind energy. However, for a given location, the cheapest H_2 is obtained from a hybrid solar and wind system. In terms of the storage of H_2 , tanks are more expensive than salt caverns. From a location perspective, the cheapest H_2 would be produced in GB.



than those obtained from the latter, with externalities (*i.e.*, 6–18% cheaper) and without externalities (*i.e.*, 4–13% cheaper).

As depicted in Fig. 6, e-jet fuels are much more expensive than fossil jet fuel (BAU), *i.e.*, 5.4–15.1-fold more costly, even

after accounting for externalities, *i.e.*, 2.3–6.8-fold more expensive. The current NPC of fossil jet fuel is 0.17 and 0.49 USD per t per km, without and with externalities, respectively.

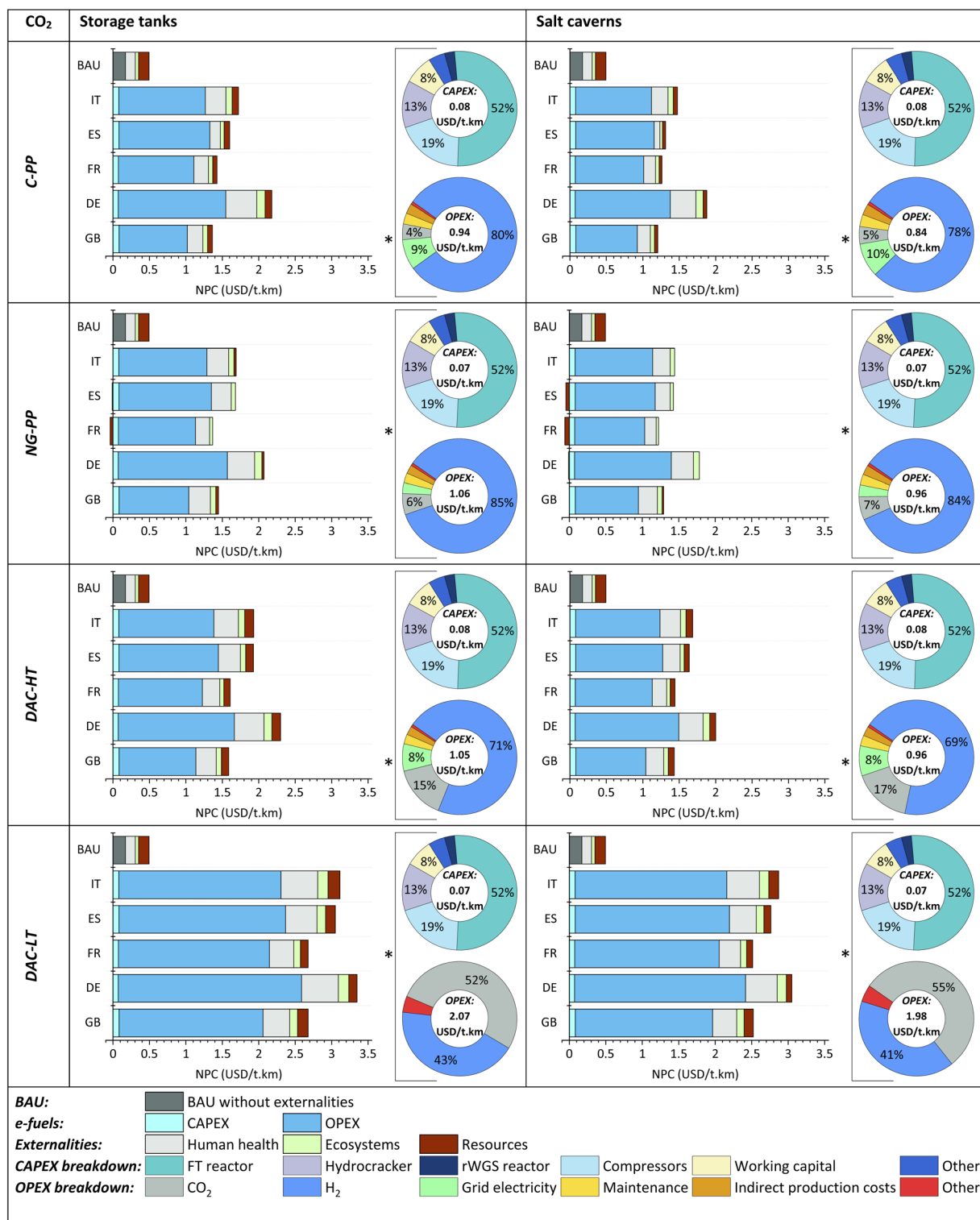


Fig. 6 e-Jet fuel: NPC breakdown. e-Jet fuels are currently more expensive than their fossil analogues even when externalities are included in the production cost. The main contributions to the NPC of e-jet fuels come from H₂, CO₂ and grid electricity. The cheapest e-jet fuel would be produced in GB using CO₂ from coal-fired plants and H₂ from hybrid solar and wind energy.



Based on the economic performance excluding externalities, the option from (C-PP)CO₂ in GB has the lowest production cost, with values of 0.93 and 1.02 USD per t per km, for H₂ stored in salt caverns and type I tanks, respectively. However, after accounting for externalities, the alternative from (NG-PP)CO₂ in FR is the cheapest option, with values of 1.16 and 1.33 USD per t per km for H₂ stored in caverns and tanks, respectively. For each of the CO₂ sources, e-jet fuel production in DE is the most costly among the locations assessed, including and excluding externalities.

OPEX is substantially higher than CAPEX in all cases, *i.e.*, 11.5–34.8-fold higher for H₂ tank storage and 10.3–32.4-fold higher for salt cavern storage.

In terms of the CAPEX of the plant, the FT reactor is the major contributor, followed by the compressors and the hydrocracker. Meanwhile, the most significant contributor to the OPEX is H₂ and, to a lesser extent, CO₂ and grid electricity, except for (DAC-LT) e-jet fuel, where CO₂ is the main contributor to the NPC, followed by H₂. It is important to mention that the CO₂ cost for DAC-LT is based on the only DAC installation currently generating negative emissions, which is located in Iceland.⁸⁵

According to the NPC breakdown of e-jet fuels, H₂ and CO₂ are the main cost contributors; their shares range between 34% and 84% for H₂ without externalities (and 26–70% with externalities) and between 3% and 55% for CO₂ without externalities (and 2–43%, with externalities).

The cost of the fuel represents only 17.7% of the total operating cost of a passenger airline,⁸⁶ as shown in Fig. 7. However, the prohibitively high production costs of e-jet fuels, even when

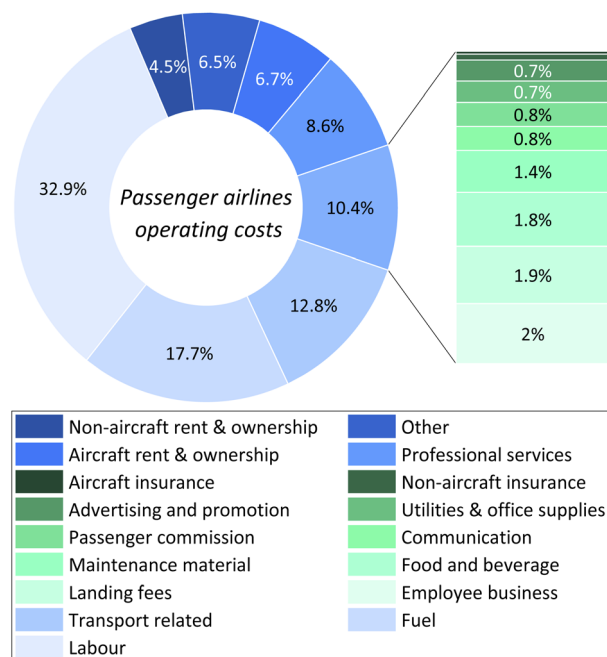


Fig. 7 Breakdown of the operating costs of passenger airlines. Approximately, 17.7% of the total operating costs of passenger airlines comes from fuel cost. However, the major contributor to these costs is labour cost, which accounts for ca. 32.9% of them.⁸⁶

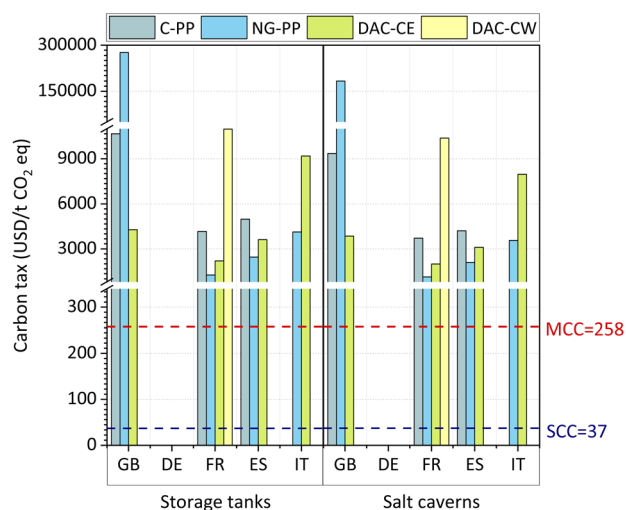


Fig. 8 Carbon tax. A minimum carbon tax of 1125 USD/tCO₂-eq would be required to make e-jet economically appealing compared to fossil jet fuel.

externalities are included, indicate that widespread deployment of these fuels will not occur in the short term, given the drastic reductions in captured CO₂ and green H₂ costs that would be necessary to make them economically appealing.

3.3.2. Abatement cost of carbon emissions. As depicted in Fig. 8, the abatement costs vary according to the CO₂ source and H₂ storage option, in the range of 1125–275 770 USD/tCO₂-eq, excluding externalities. All these values lie well above the current estimated social cost of carbon emissions (SCC), *i.e.*, 37 USD/tCO₂-eq, and mortality cost of carbon emissions (MCC), *i.e.*, 258 USD/tCO₂-eq.⁸⁷

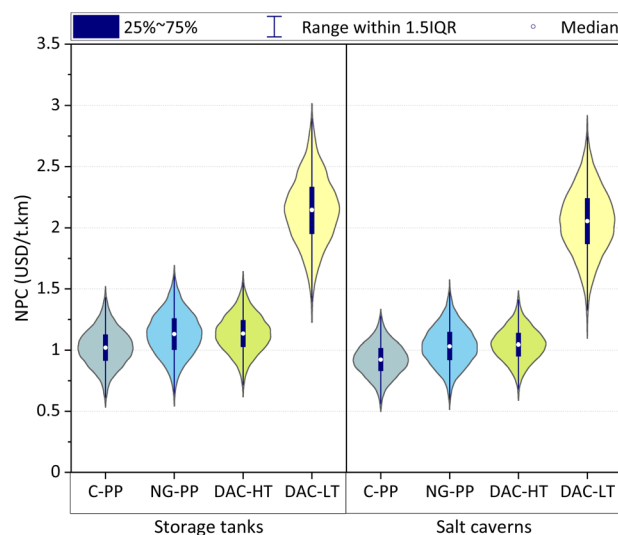


Fig. 9 Uncertainty analysis. Based on the given parameters, the NPC of e-jet can go as low as 0.46/0.36 USD per t per km (from (C-PP) CO₂), 0.53/0.39 USD per t per km (from (NG-PP) CO₂), 0.52 USD per t per km (from (DAC-HT) CO₂), and 1.11/1.03 USD per t per km (from (DAC-LT) CO₂), for tanks and salt caverns storage, respectively.



3.3.3. Uncertainty analysis. The results of the Monte Carlo simulations are shown in Fig. 9.

The variance of the cost lies in the range of 0.46–3.27 and 0.36–2.96 USD per t per km for H₂ stored in tanks and salt caverns, respectively, indicating that the NPC of e-jet fuels has a strong dependence on the cost of H₂, CO₂ and grid electricity.

Regardless of the H₂ storage option, e-jet fuel from (C-PP) CO₂ is the cheapest alternative on average (1.02 and 0.92 USD per t per km for storage in tanks and caverns, respectively), followed by e-jet fuel produced using CO₂ from NG-PP (1.13 and 1.03 USD per t per km), DAC-HT (1.13 and 1.04 USD per t per km) and DAC-LT (2.14 and 2.05 USD per t per km).

4. Conclusions

This work presents a critical analysis of e-jet fuel production from different sources of CO₂ and green H₂, using a wide range of tools to evaluate its current economic and environmental performance and elucidate the potential challenges for its widespread utilisation. Based on our analysis, e-jet fuel is not economically competitive against fossil fuel (*i.e.*, at least 5.4-fold the cost of the fossil counterpart), mainly owing to the high price of green H₂ and captured CO₂, the two main feedstocks. This holds true even when the monetised LCA endpoint impacts, *i.e.*, externalities, are added to the production cost; however, the economic gap is reduced such that e-jet fuel is found to be approximately 2.3-fold more expensive than the fossil analogue.

For e-jet fuel to be economically competitive, a tax on CO₂ of at least 1125 USD/t_{CO₂-eq} should be imposed on the fossil counterpart. This tax is equivalent to 30.4-fold the social cost of carbon and 4.4-fold the mortality cost of carbon at present. Thus, this is not deemed realistic.

From the environmental perspective, e-jet fuels could exacerbate some environmental categories. Whilst e-jet fuels mostly outperform fossil jet fuels in terms of global warming potential and resource scarcity, human health and ecosystem quality are worsened, *i.e.*, burden-shifting from resource scarcity towards human health and ecosystem quality.

In view of the impact that CO₂ and H₂ have on the environmental and economic performance of e-jet fuels, efforts should focus on reducing their cost and impact, *i.e.*, CO₂ capture and H₂O electrolysis, respectively, especially with regard to the electricity used, which is the most significant contributor to costs and environmental impacts (high carbon intensity at the moment).

Salt caverns for H₂ storage and CO₂ from natural gas and coal-fired power plants have proven to be among the cheapest alternatives; however, salt caverns are not universally available, and the use of fossil CO₂ as a feedstock is fundamentally unsustainable. Hence, except for particular cases, H₂ storage in tanks and CO₂ from DAC will be considered the standard for e-jet fuel production plants requiring constant output.

As CO₂ is also generated from burning e-jet fuels, where fossil CO₂ is used as a feedstock, this technology can only be seen as an intermediate step towards a net-zero economy. In this regard, carbon capture and storage/sequestration (CCS)

and bioenergy with carbon capture and storage/sequestration (BECCS) options should be evaluated as a possible way to close the carbon loop and effectively remove CO₂ from the atmosphere.

The analysis presented in this study could be extended to regions other than Europe. The production costs of e-fuels would vary, depending mainly on the available energy resources (*i.e.*, solar radiation and wind speed)^{34,35} and the investment risk associated with each region.^{88,89} Notwithstanding this, e-fuels are not expected to be economically competitive against fossil fuels using current technology. This is mainly due to the high investment costs of green H₂ and, to a lesser extent, captured CO₂ – the main raw materials for producing e-fuels.

Overall, the results of our study indicate that if we have the potential to deploy decarbonised energy, then it is better used to displace higher-carbon assets from the grid rather than to produce e-fuels. This work also highlights the importance of assessing any new technology oriented to sustainability beyond traditional economic and global warming potential indicators, given the potential for environmental burden shifting. Comprehensive assessments encompassing the economic performance and the global overview of the potential environmental impacts posed by any new technology will provide a more comprehensive picture that will facilitate more informed decision-making. Furthermore, we showed that accounting for the intermittency of renewables as part of the feasibility assessment of any project is key, given its strong impact on the project costs.

Author contributions

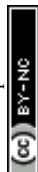
Diego Freire Ordóñez: conceptualisation, methodology, software, validation, formal analysis, investigation, data curation, writing – original draft, writing – review & editing, visualisation. Thorsteinn Halfdanarson: methodology. Caroline Ganzer: methodology. Nilay Shah: writing – review & editing, supervision. Niall Mac Dowell: writing – review & editing, supervision. Gonzalo Guillén-Gosálbez: conceptualisation, methodology, writing – review & editing, supervision.

Conflicts of interest

There are no conflicts of interest to declare.

Acknowledgements

Diego Freire has been funded by the Ecuadorian Secretariat for Higher Education, Science, Technology and Innovation, SEN-ESCYT, Award No. 106-2017. The Institute for Applied Sustainability Research, iiasur, supports international research on global sustainability applied to the Global South. This publication was also created as part of NCCR Catalysis (grant 180544), a National Centre of Competence in Research funded by the Swiss National Science Foundation.



Notes and references

- 1 H. Ritchie and M. Roser, *Energy, Our World in Data*, 2020.
- 2 J.-P. Rodrigue, *The geography of transport systems*, Routledge, Abingdon Oxon, New York NY, 2020.
- 3 Benoit Lefevre and Angela Enriquez, *Transport Sector Key to Closing the World's Emissions Gap*, 2014.
- 4 K. Dahal, S. Brynolf, C. Xisto, J. Hansson, M. Grahn, T. Grönstedt and M. Lehtveer, Techno-economic review of alternative fuels and propulsion systems for the aviation sector, *Renewable Sustainable Energy Rev.*, 2021, **151**, 111564.
- 5 I. Ridjan, B. V. Mathiesen and D. Connolly, Terminology used for renewable liquid and gaseous fuels based on the conversion of electricity, *J. Cleaner Prod.*, 2016, **112**, 3709–3720.
- 6 A. Goldmann, W. Sauter, M. Oettinger, T. Kluge, U. Schröder, J. Seume, J. Friedrichs and F. Dinkelacker, A Study on Electrofuels in Aviation, *Energies*, 2018, **11**, 392.
- 7 J. Åkerman, A. Kamb, J. Larsson and J. Nässén, Low-carbon scenarios for long-distance travel 2060, *Transportation Research Part D: Transport and Environment*, 2021, **99**, 103010.
- 8 D. Freire Ordóñez, N. Shah and G. Guillén-Gosálbez, Economic and full environmental assessment of electrofuels via electrolysis and co-electrolysis considering externalities, *Appl. Energy*, 2021, **286**, 116488.
- 9 D. Freire Ordóñez and G. Guillén-Gosálbez, in *30th european symposium on computer aided chemical engineering*, Elsevier, 2020, pp. 595–600.
- 10 D. Freire Ordóñez, T. Halfdanarson, C. Ganzer, G. Guillén-Gosálbez, N. M. Dowell and N. Shah, in *Computer Aided Chemical Engineering*, ed. A. Espuña, M. Graells and L. Puigjaner, Elsevier, 2017, pp. 1623–1628.
- 11 S. Hänggi, P. Elbert, T. Büttler, U. Cabalzar, S. Teske, C. Bach and C. Onder, A review of synthetic fuels for passenger vehicles, *Energy Reports*, 2019, **5**, 555–569.
- 12 L. Guzzella and A. Sciarretta, *Vehicle propulsion systems. Introduction to modeling and optimization*, Springer, Berlin, 3rd edn, 2013.
- 13 M. Lehtveer, S. Brynolf and M. Grahn, What Future for Electrofuels in Transport? Analysis of Cost Competitiveness in Global Climate Mitigation, *Environ. Sci. Technol.*, 2019, **53**, 1690–1697.
- 14 S. J. Davis, N. S. Lewis, M. Shaner, S. Aggarwal, D. Arent, I. L. Azevedo, S. M. Benson, T. Bradley, J. Brouwer, Y.-M. Chiang, C. T. M. Clack, A. Cohen, S. Doig, J. Edmonds, P. Fennell, C. B. Field, B. Hannegan, B.-M. Hodge, M. I. Hoffert, E. Ingersoll, P. Jaramillo, K. S. Lackner, K. J. Mach, M. Mastrandrea, J. Ogden, P. F. Peterson, D. L. Sanchez, D. Sperling, J. Stagner, J. E. Trancik, C.-J. Yang and K. Caldeira, Net-zero emissions energy systems, *Science*, 2018, **360**, eaas9793.
- 15 M. M. Alhyari, A. Al-Salaymeh, M. R. Irshidat, M. Kaltschmitt and U. Neuling, The Impact of Energy Source on the Life-Cycle Assessment of Power-to-Liquid Fuels, *Journal of Ecological Engineering*, 2019, **20**, 239–244.
- 16 K. Kieckhäfer, G. Quante, C. Müller, T. Spengler, M. Lossau and W. Jonas, Simulation-Based Analysis of the Potential of Alternative Fuels towards Reducing CO₂ Emissions from Aviation, *Energies*, 2018, **11**, 186.
- 17 S. Schemme, J. L. Breuer, M. Köller, S. Meschede, F. Walman, R. C. Samsun, R. Peters and D. Stolten, H₂-based synthetic fuels: A techno-economic comparison of alcohol, ether and hydrocarbon production, *Int. J. Hydrogen Energy*, 2020, **45**(8), 5395–5414.
- 18 P. Schmidt, V. Batteiger, A. Roth, W. Weindorf and T. Raksha, *Power to Liquids as Renewable Fuel Option for Aviation. A Review*, 2018.
- 19 P. Schmidt, W. Weindorf, A. Roth, V. Batteiger, F. Riegel and Deutschland Umweltbundesamt, *Power-to-liquids: Potentials and Perspectives for the Future Supply of Renewable Aviation Fuel*, German Environment Agency, 2016.
- 20 E. D. Sherwin, Electrofuel Synthesis from Variable Renewable Electricity: An Optimization-Based Techno-Economic Analysis, *Environ. Sci. Technol.*, 2021, **55**, 7583–7594.
- 21 J. Trinh, F. Harahap, A. Fagerström and J. Hansson, What Are the Policy Impacts on Renewable Jet Fuel in Sweden?, *Energies*, 2021, **14**, 7194.
- 22 G. Hutchings, M. Davidson, P. Atkins, P. Collier, N. Jackson, A. Morton, M. Muskett, M. Rosseinsky, P. Styring, P. Thornley and C. Williams, *Sustainable synthetic carbon based fuels for transport. Sustainable synthetic carbon based fuels for transport*, The Royal Society, London, 2019.
- 23 J. Armijo and C. Philibert, Flexible production of green hydrogen and ammonia from variable solar and wind energy: Case study of Chile and Argentina, *Int. J. Hydrogen Energy*, 2020, **45**, 1541–1558.
- 24 C. Ganzer and N. Mac Dowell, A comparative assessment framework for sustainable production of fuels and chemicals explicitly accounting for intermittency, *Sustainable Energy Fuels*, 2020, **4**(8), 3888–3903.
- 25 B. P. Weidema, Comparing Three Life Cycle Impact Assessment Methods from an Endpoint Perspective, *J. Ind. Ecol.*, 2015, **19**, 20–26.
- 26 International Energy Agency (IEA) Data, International Energy Agency.
- 27 Aspen Technology, *Aspen Plus*, <https://www.aspentech.com/en/products/engineering/aspen-plus>.
- 28 D. H. König, M. Freiberg, R.-U. Dietrich and A. Wörner, Techno-economic study of the storage of fluctuating renewable energy in liquid hydrocarbons, *Fuel*, 2015, **159**, 289–297.
- 29 F. G. Albrecht, D. H. König, N. Baucks and R.-U. Dietrich, A standardized methodology for the techno-economic evaluation of alternative fuels – a case study, *Fuel*, 2017, **194**, 511–526.
- 30 O. Schmidt, A. Gambhir, I. Staffell, A. Hawkes, J. Nelson and S. Few, Future cost and performance of water electrolysis: An expert elicitation study, *Int. J. Hydrogen Energy*, 2017, **42**, 30470–30492.
- 31 PV magazine, *Portuguese consortium plans 1 GW green hydrogen cluster*, <https://www.pv-magazine.com/2020/07/30/>



- portuguese-consortium-plans-1-gw-green-hydrogen-cluster/, accessed 7 January 2022.
- 32 PV magazine, *Thyssenkrupp increases annual electrolyzer capacity to 1 GW*, <https://www.pv-magazine.com/2020/06/09/thyssenkrupp-increases-annual-electrolyzer-capacity-to-1-gw/>, accessed 7 January 2022.
 - 33 Recharge, *Plans unveiled for 1GW green-hydrogen power plant fuelled by wind and solar | Recharge*, <https://www.rechargenews.com/transition/plans-unveiled-for-1gw-green-hydrogen-power-plant-fuelled-by-wind-and-solar/2-1-812928>, accessed 7 January 2022.
 - 34 S. Pfenninger and I. Staffell, Long-term patterns of European PV output using 30 years of validated hourly reanalysis and satellite data, *Energy*, 2016, **114**, 1251–1265.
 - 35 I. Staffell and S. Pfenninger, Using bias-corrected reanalysis to simulate current and future wind power output, *Energy*, 2016, **114**, 1224–1239.
 - 36 G. Reiter and J. Lindorfer, Evaluating CO₂ sources for power-to-gas applications – A case study for Austria, *J. CO₂ Util.*, 2015, **10**, 40–49.
 - 37 S. Sengupta, A. Jha, P. Shende, R. Maskara and A. K. Das, Catalytic performance of Co and Ni doped Fe-based catalysts for the hydrogenation of CO₂ to CO via reverse water-gas shift reaction, *J. Environ. Chem. Eng.*, 2019, **7**, 102911.
 - 38 R. B. Unde, *Kinetics and reaction engineering aspects of syngas production by the heterogeneously catalysed reverse water gas shift reaction*, Germany, 2012.
 - 39 D. H. König, N. Baucks, R.-U. Dietrich and A. Wörner, Simulation and evaluation of a process concept for the generation of synthetic fuel from CO₂ and H₂, *Energy*, 2015, **91**, 833–841.
 - 40 Z. Zhang, C. Shen, K. Sun, X. Jia, J. Ye and C. Liu, Advances in studies of the structural effects of supported Ni catalysts for CO₂ hydrogenation: from nanoparticle to single atom catalyst, *J. Mater. Chem. A*, 2022, **10**, 5792–5812.
 - 41 M. Hillestad, Modeling the Fischer-Tropsch Product Distribution and Model Implementation, *Chem. Prod. Process Model.*, 2015, **10**, 147–159.
 - 42 M. Samavati, *Design and analysis of solid oxide electrolysis-based systems for synthetic liquid fuels production*, KTH Royal Institute of Technology, 2018.
 - 43 D. Selvatico, A. Lanzini and M. Santarelli, Low Temperature Fischer-Tropsch fuels from syngas: Kinetic modeling and process simulation of different plant configurations, *Fuel*, 2016, **186**, 544–560.
 - 44 N. Jungbluth, R. Frischknecht, M. Faist Emmenegger, R. Steiner and M. Tuchscheid, *Life Cycle Assessment of BTL-fuel production: Inventory Analysis*, 2007.
 - 45 K. Ibrik, *Kinetics of the Fischer-Tropsch Reaction Over Alumina Supported Cobalt Catalyst in a Slurry Reactor*, Qatar Foundation Annual Research Forum Proceedings, 2011, EGPS2.
 - 46 D. Vervloet, F. Kapteijn, J. Nijenhuis and J. R. van Ommen, Fischer-Tropsch reaction-diffusion in a cobalt catalyst particle: aspects of activity and selectivity for a variable chain growth probability, *Catal. Sci. Technol.*, 2012, **2**, 1221.
 - 47 D. H. König, M. Freiberg, R.-U. Dietrich and A. Wörner, Techno-economic study of the storage of fluctuating renewable energy in liquid hydrocarbons, *Fuel*, 2015, **159**, 289–297.
 - 48 M. Moser, T. Pregger, S. Simon, D. H. König, A. Wörner, R. U. Dietrich, M. Köhler, P. Oßwald, J. Grohmann, T. Kathrotia, G. Eckel, D. Schweitzer, N. Armbrust, H. Dieter, G. Scheffknecht, C. Kern, J. Thiessen, A. Jess and M. Aigner, Synthetic Liquid Hydrocarbons from Renewable Energy – Results of the Helmholtz Energy Alliance, *Chem.-Ing.-Tech.*, 2017, **89**, 274–288.
 - 49 B. Viswanathan, in *Energy sources*, ed. B. Viswanathan, Elsevier, Amsterdam, Boston, 2017, pp. 29–57.
 - 50 P. R. Robinson, in *Advances in Clean Hydrocarbon Fuel Processing: Woodhead Publishing Series in Energy*, ed. M. R. Khan, Woodhead Publishing, 2011, pp. 287–325.
 - 51 J. Lee, S. Hwang, J. G. Seo, S.-B. Lee, J. C. Jung and I. K. Song, Production of middle distillate through hydrocracking of paraffin wax over Pd/SiO₂-Al₂O₃ catalysts, *J. Ind. Eng. Chem.*, 2010, **16**, 790–794.
 - 52 J. Lee, S. Hwang, J. G. Seo, U. G. Hong, J. C. Jung and I. K. Song, Pd catalyst supported on SiO₂-Al₂O₃ xerogel for hydrocracking of paraffin wax to middle distillate, *J. Ind. Eng. Chem.*, 2011, **17**, 310–315.
 - 53 Y. Liu, K. Murata, K. Okabe, M. Inaba, I. Takahara, T. Hanaoka and K. Sakanishi, Selective hydrocracking of Fischer-Tropsch waxes to high-quality diesel fuel over pt-promoted polyoxocation-pillared montmorillonites, *Top. Catal.*, 2009, **52**, 597–608.
 - 54 V. Calemme, C. Gambaro, W. O. Parker, R. Carbone, R. Giardino and P. Scorletti, Middle distillates from hydrocracking of FT waxes: composition, characteristics and emission properties, *Catal. Today*, 2010, **149**, 40–46.
 - 55 T. Hengsawad, C. Srimingkwanchai, S. Butnark, D. E. Resasco and S. Jongpatiwut, Effect of metal-acid balance on hydroprocessed renewable jet fuel synthesis from hydrocracking and hydroisomerization of biohydrogenated diesel over Pt-supported catalysts, *Ind. Eng. Chem. Res.*, 2018, **57**, 1429–1440.
 - 56 M. M. El-Halwagi, *Sustainable Design Through Process Integration: Fundamentals and Applications to Industrial Pollution Prevention, Resource Conservation, and Profitability Enhancement*, Elsevier Ltd, Amsterdam, 2nd edn, 2017.
 - 57 Aspen Technology, *Aspen Energy Analyzer*, <https://www.aspentech.com/en/products/pages/aspen-energy-analyzer>.
 - 58 H. Bruijn, R. Duin, M. A. J. Huijbregts, J. B. Guinee, M. Gorree, R. Heijungs, G. Huppes, R. Kleijn, A. Koning, L. Oers and A. Wegener Sleeswijk, *Handbook on Life Cycle Assessment. Operational Guide to the ISO Standards*, Kluwer Academic Publishers, Dordrecht, 2004.
 - 59 Pré Consultants B. V., *SimaPro|The World's Leading LCA Software*, <https://simapro.com/>, (accessed 2 August 2019).
 - 60 L. J. Müller, A. Kätelhön, M. Bachmann, A. Zimmermann, A. Sternberg and A. Bardow, A Guideline for Life Cycle Assessment of Carbon Capture and Utilization, *Front. Energy Res.*, 2020, **8**, 15.



- 61 K. Bareiß, C. de La Rua, M. Möckl and T. Hamacher, Life cycle assessment of hydrogen from proton exchange membrane water electrolysis in future energy systems, *Appl. Energy*, 2019, **237**, 862–872.
- 62 S. Häfele, M. Hauck and J. Dailly, Life cycle assessment of the manufacture and operation of solid oxide electrolyser components and stacks, *Int. J. Hydrogen Energy*, 2016, **41**, 13786–13796.
- 63 B. Sørensen and G. Spazzafumo, in *Hydrogen and Fuel Cells*, ed. B. Sørensen and G. Spazzafumo, Elsevier Science, San Diego, CA, USA, 2018, pp. 413–461.
- 64 H. Nojoumi, I. Dincer and G. F. Naterer, Greenhouse gas emissions assessment of hydrogen and kerosene-fueled aircraft propulsion, *Int. J. Hydrogen Energy*, 2009, **34**, 1363–1369.
- 65 G. Wernet, C. Bauer, B. Steubing, J. Reinhard, E. Moreno-Ruiz and B. Weidema, The ecoinvent database version 3 (part I): overview and methodology, *Int. J. Life Cycle Assess.*, 2016, **21**, 1218–1230.
- 66 M. Tomatis, M. T. Moreira, H. Xu, W. Deng, J. He and A. M. Parvez, Removal of VOCs from waste gases using various thermal oxidizers: A comparative study based on life cycle assessment and cost analysis in China, *J. Cleaner Prod.*, 2019, **233**, 808–818.
- 67 M. A. J. Huijbregts, Z. J. N. Steinmann, P. M. F. Elshout, G. Stam, F. Veronesi, M. Vieira, M. Zijp, A. Hollander and R. van Zelm, ReCiPe2016: a harmonised life cycle impact assessment method at midpoint and endpoint level, *Int. J. Life Cycle Assess.*, 2017, **22**, 138–147.
- 68 R. Sinnott and G. Towler, *Chemical engineering design*, Elsevier, Oxford, 2020.
- 69 L. L. Access Intelligence, *The Chemical Engineering Plant Cost Index - Chemical Engineering*, <https://www.chemengonline.com/pci-home>, (accessed 18 April 2020).
- 70 World Bank, *Commodity Markets*, <https://www.worldbank.org/en/research/commodity-markets>, (accessed 11 January 2022).
- 71 M. B. Viswanathan, D. R. Raman, K. A. Rosentrater and B. H. Shanks, A Technoeconomic Platform for Early-Stage Process Design and Cost Estimation of Joint Fermentative–Catalytic Bioprocessing, *Processes*, 2020, **8**, 229.
- 72 IATA, *Economic Performance of the Airline Industry*, 2020.
- 73 B. Graver, K. Zhang and D. Rutherford, *CO₂ emissions from commercial aviation*, 2018, 2022, <https://theicct.org/publications/co2-emissions-commercial-aviation-2018>, (accessed 11 January 2022).
- 74 Laia and Pierre-Selim, *How much fuel per passenger an aircraft is consuming?*, <https://blog.openairlines.com/how-much-fuel-per-passenger-an-aircraft-is-consuming>, (accessed 11 January 2022).
- 75 D. F. Rodríguez-Vallejo, G. Guillén-Gosálbez and B. Chachuat, What Is the True Cost of Producing Propylene from Methanol? The Role of Externalities, *ACS Sustainable Chem. Eng.*, 2020, **8**, 3072–3081.
- 76 I. M. Algunaibet, C. Pozo, Á. Galán-Martín and G. Guillén-Gosálbez, Quantifying the cost of leaving the Paris Agreement via the integration of life cycle assessment, energy systems modeling and monetization, *Appl. Energy*, 2019, **242**, 588–601.
- 77 A. González-Garay, M. S. Frei, A. Al-Qahtani, C. Mondelli, G. Guillén-Gosálbez and J. Pérez-Ramírez, Plant-to-planet analysis of CO₂-based methanol processes, *Energy Environ. Sci.*, 2019, **12**, 3425–3436.
- 78 O. Schmidt, S. Melchior, A. Hawkes and I. Staffell, Projecting the Future Levelized Cost of Electricity Storage Technologies, *Joule*, 2019, **3**, 81–100.
- 79 M. Pehl, A. Arvesen, F. Humpenöder, A. Popp, E. G. Hertwich and G. Luderer, Understanding future emissions from low-carbon power systems by integration of life-cycle assessment and integrated energy modelling, *Nat. Energy*, 2017, **2**, 939–945.
- 80 D. G. Caglayan, N. Weber, H. U. Heinrichs, J. Linßen, M. Robinius, P. A. Kukla and D. Stolten, Technical potential of salt caverns for hydrogen storage in Europe, *Int. J. Hydrogen Energy*, 2020, **45**, 6793–6805.
- 81 Fuels Cells and Hydrogen 2 Joint Undertaking, *Opportunities for Hydrogen Energy Technologies Considering the National Energy & Climate Plans*, 2020.
- 82 D. W. Keith, G. Holmes, D. St. Angelo and K. Heidel, A Process for Capturing CO₂ from the Atmosphere, *Joule*, 2018, **2**, 1573–1594.
- 83 S. Deutz and A. Bardow, Life-cycle assessment of an industrial direct air capture process based on temperature–vacuum swing adsorption, *Nat. Energy*, 2021, **6**, 203–213.
- 84 N. Mac Dowell, N. Sunny, N. Brandon, H. Herzog, A. Y. Ku, W. Maas, A. Ramirez, D. M. Reiner, G. N. Sant and N. Shah, The hydrogen economy: A pragmatic path forward, *Joule*, 2021, **5**, 2524–2529.
- 85 H. J. Herzog, *Direct air capture: A process engineer's view*, 2021.
- 86 Airlines For America, *A4A Passenger Airline Cost Index (PACI)*, <https://www.airlines.org/dataset/a4a-quarterly-passenger-airline-cost-index-u-s-passenger-airlines/>.
- 87 R. D. Bressler, The mortality cost of carbon, *Nat. Commun.*, 2021, **12**, 1–12.
- 88 B. U. Schyska and A. Kies, How regional differences in cost of capital influence the optimal design of power systems, *Appl. Energy*, 2020, **262**, 114523.
- 89 C. Agutu, F. Egli, N. J. Williams, T. S. Schmidt and B. Steffen, Accounting for finance in electrification models for sub-Saharan Africa, *Nat. Energy*, 2022, 1–11.

