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From fossil to green chemicals: sustainable pathways and new carbon feedstocks for the global chemical industry[†]

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Following current trends, the global chemical industry is set to become the largest consumer of fossil fuels. Among energy intensive industries, the chemical industry is one of the most challenging to defossilise due to the abundance of cheap fossil fuel-feedstocks and it is currently responsible for roughly 3% of global anthropogenic CO₂ emissions. Unlike other energy-intensive industries, the chemical industry cannot be made fully sustainable directly with renewable electricity and green electricity-based hydrogen (e-hydrogen). Therefore, new green carbon feedstocks must be developed to defossilise the production of large volume organic chemicals. The most promising green carbon feedstocks are electricity-based methanol (e-methanol) and biomass-based methanol (bio-methanol), which can be used directly or as a feedstock for olefin and aromatic production. Increased recycling of plastics will reduce the amount of primary feedstock that will be required for chemical production. To investigate the energy and feedstock requirements for a global defossilisation of chemical production, scenarios are developed that reach net-zero emissions by 2040, 2050, and 2060 compared to business-as-usual conditions to 2100. High and low biomass feedstock variations are included to investigate the potential of biomass feedstocks in the future chemical industry, which are limited due to strict sustainability criteria. The results suggest that the chemical industry could become the largest e-hydrogen consumer, with a demand ranging from 16100 to 23100 TWh_{H21HV} in 2050. High shares of electricity-based chemicals (e-chemicals) were found to provide the lowest annualised costs, suggesting that an e-chemical transition pathway may be the most economically competitive pathway to defossilise the global chemical industry.

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Broader context

While the defossilisation of energy systems is well understood, the decoupling of fossil fuels from the chemical industry has often been overlooked, due to the abundance of cheap fossil feedstocks, which are used to produce plastics, fertilisers, pesticides, fibres, and personal care and consumer products, among other ubiquitous chemicals. The key challenge with the defossilisation of the chemical industry is the requirement of carbon-based feedstocks, which cannot be directly substituted with renewable electricity. Chemical demand is also expected to grow rapidly in the coming decades, and, without major disruption to feedstocks, could become the largest driver in oil consumption. Renewable electricity- and biomass-based feedstocks have been suggested to substitute fossil feedstocks; however, there is a knowledge gap in the energy system requirements to completely replace fossil feedstocks. This research presents scenarios for the complete defossilisation of global chemical feedstocks from 2020 to 2100 using a high geographical resolution of 145 regions. The results of this study found that the complete defossilisation of chemical production applying high levels of power-to-chemicals, in tandem with increased plastic recycling, will lead to the lowest annualised costs by 2050.

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1. Introduction

1.1. Current situation and technologies

Chemical products have become ubiquitous in modern society and are integral to the functioning of modern society. The demand for chemical products, especially plastics, has grown faster than that for any other bulk material.¹ From 1950 to



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2015, resins and fibres used in plastic production have grown at a compound annual growth rate (CAGR) of 8.4%, which has been estimated to be around 2.5 times higher than the CAGR of the global gross domestic product.² Despite being the largest industrial energy consumer of both oil and gas, the chemical industry was only the third largest industrial emitter at 920 MtCO2 in 2019 resulting from primary chemical production alone, behind the cement, and iron and steel industries.³ This is largely due to the high shares of non-energy use of fossil fuels as feedstocks where high shares of the fossil carbon are embedded in the chemicals produced. Following current trends, demand for chemical products in the form of fertilisers, pesticides, plastics, and fibres, among others, is expected to increase significantly in the coming decades,^{1,2,4,5} and estimates suggest that, without intervention, non-combusted feedstocks will become the largest source of fossil fuel demand growth. The use of chemical products will additionally be essential to produce technologies that can eliminate emissions in the energy sector.⁶ Therefore, to meet the targets of the Paris Agreement⁷ and develop a carbon-neutral and sustainable chemical industry, a complete defossilisation of carbon feedstocks must occur.

The foundations of the modern organic chemical industry are built on seven key building blocks or primary chemicals: ammonia (NH₃), methanol (CH₃OH or MeOH), ethylene (C₂H₄), propylene (C₃H₆), benzene (C₆H₆), toluene (C₇H₈), and mixed xylenes (C₈H₁₀). Ethylene and propylene are often discussed as light olefins, and benzene, toluene, and mixed xylenes are referred to as BTX aromatics, and together are referred to as high value chemicals (HVCs). Due to the abundance of low-cost fossil feedstocks, discussions of emission reductions in the chemical industry have largely focused on process emissions.¹ However, many chemical processes have reached close to their highest feedstock efficiency; however, processes may be improved to reduce CO_2 emissions.^{1,8} Today, primary chemical production has emission factors of 2.4 t CO_2 /tNH₃, 2.3 t CO_2 /tMeOH, and 1.0 t CO_2 /tHVC,¹ related to both the fossil feedstocks required and process heat and electricity.

The conventional production of ammonia uses the Haber-Bosch process with fossil methane, coal, or oil as the feedstock. as shown in Fig. 1, and fossil methane has become the dominant feedstock globally, as it is responsible for 75% of global ammonia feedstocks, followed by coal at 22% and oil at 3%.8 With fossil feedstocks, this process is responsible for around 1.8% of global CO₂ emissions.⁹ Ammonia production has grown exponentially since the development of the Haber-Bosch process in 1909, and its global production reached 195 MtNH₃ in 2018.¹⁰ In addition to its use as a fertiliser, ammonia has been discussed as a form of seasonal storage to offset variable renewable energy (RE) and as a fuel for the transport sector, especially in marine applications.¹¹⁻¹⁷ From both a feedstock perspective and an emission perspective, the fossil methane-to-ammonia route has the best performance, with a feedstock requirement of 28 $\text{GJ}_{\text{CH4,LHV}}/\text{tNH}_3^{-13,18}$ and an emission factor of 1.6 tCO₂/tNH₃.¹⁸ However, as highlighted by Smith et al.,11 only marginal efficiency improvements are available for conventional ammonia production.

Methanol production, as shown in Fig. 2, similarly converts syngas, a mixture of H_2 , CO, and CO₂, from a fossil fuel feedstock to methanol¹⁹ using a Cu/ZnO/Al₂O₃ catalyst.²⁰ Similar to the ammonia production process, steam methane reformation is used for fossil methane and naphtha feedstocks, whereas partial oxidation is used for heavy oils and solid fossil fuels. Fossil methane is the most used feedstock for methanol



Fig. 1 Conventional production route for ammonia production from natural gas and nitrogen from an air separation unit (ASU). Coal and oil can similarly be used as a fossil feedstock, using coal gasification and partial oxidation for syngas production, respectively. Before nitrogen and hydrogen are input to the Haber–Bosch reactor, an additional methanation step is required to convert carbon monoxide and carbon dioxide to methane that accumulates as inert in the NH₃ synthesis stage, minimising the poisoning of the Haber–Bosch catalyst. The condenser after the Haber–Bosch reactor removes H₂ and N₂ impurities from the outlet stream. Adapted from Smith *et al.*¹¹



synthesis, corresponding to 57% of global methanol feedstocks,⁸ and has a feedstock demand of 33.9 GJ_{CH4,LHV}/tMeOH.²¹ Coal is the next most used feedstock at 40% of global methanol feedstocks consuming 46.9 GJ_{CH4,LHV}/tMeOH,²¹ largely due to the high shares of methanol production and consumption in China,¹⁹ and oil composes only 3% of global methanol feedstocks, for a total production of 95 MtMeOH in 2018.¹⁰ Additionally, CO₂ emissions for the fossil-based methanol synthesis range from 0.5 tCO_{2eq}/tMeOH for steam reforming to 1.5 tCO_{2eq}/tMeOH for partial oxidation.²²

HVCs are very often co-produced in several chemical processes or produced as by-products from refineries. The most widely used process for the co-production of ethylene, propylene, and BTX aromatics is the steam cracker, and is shown in Fig. 3. While naphtha is used as the feedstock, there are a wide range of feedstocks that are used for steam crackers, which largely varies on what low-cost fossil feedstock is readily available regionally. In Europe and Eurasia, heavier oil feedstocks including naphtha and gas oil are used due to their availability compared to lighter natural gas liquid feedstocks such as ethane and propane being preferred in North America and the Middle East and North Africa (MENA) regions.¹⁰ The composition of the feedstock has a significant effect on the shares of products produced, and ethylene and propylene yields from various feedstocks and operating conditions can range from 24 to 55% and 1.5 to 18%, respectively.²³ Along with propylene from steam cracking,

propylene is also produced in large quantities *via* refinery operations through deep catalytic cracking (DCC) as well as *via* propane dehydrogenation, a form of on-purpose propylene (OPP).²⁴ The global production of ethylene and propylene reached 160 and 107 Mt in 2018, respectively.¹⁰

For heavier steam cracker feedstocks and DCC, as shown in Fig. 4, BTX aromatics are produced in a pyrolysis gas (pygas) component.

However, the majority of the BTX aromatics are sourced *via* the catalytic reforming of naphtha (CRR) in refineries,^{1,8} shown in Fig. 5. Additionally, toluene can be converted to benzene or mixed xylenes through toluene hydrodealkylation (THD) and disproportionation (TDP), which has been used to reduce the overproduction of toluene relative to demand.¹⁰ In 2018, the global production of BTX aromatics reached 43, 22, and 54 Mt for benzene, toluene, and mixed xylenes (*ortho/meta/para*), respectively.¹⁰

Of the global chemical flows, roughly 61% of downstream chemicals considered in this study are used for plastic production, and comprise roughly 24% of all global chemical flows.⁸ Therefore, increasing plastic recycling has become a target of many industrialised economies that will reduce the primary chemical demand, particularly for HVCs.¹ Global plastic collection rates for plastics have been increasing steadily since 1990, increasing from 2% to 15% in 2019.²⁸ However, there is significant disparity between plastic recycling rates by regions,



Fig. 3 Conventional steam cracking process to produce ethylene, propylene, and BTX aromatics from naphtha. The pyrolysis furnace converts the feedstock, here, naphtha, into cracking gas, which is then sent to a series of separation units to separate the various hydrocarbon fractions and waste water. Additional feedstocks for the steam cracking process include ethane, propane, butane, and gas oil. Adapted from Xiang *et al.*²⁵

ranging from 8.2% in non-OECD other Africa to 25.2% in OECD Europe for 2019, as shown in Fig. $6.^{28}$

Additionally, the actual recycling rate of collected plastics is still rather low globally, ranging from 50% in the United States to 71% in non-OECD Latin America.²⁸ Although plastic recycling rates have grown rather significantly, global secondary plastic production has not seen the same levels of growth, as secondary plastic production has only grown from 1.5% of total plastics in 1990 to 6.3% in 2019, as shown in Fig. 7.

Unlike other energy-intensive materials, plastics tend to have short lifetimes, Table 1 shows the main uses of plastics, shares of plastics by use, and mean product lifetimes according to Geyer *et al.*,² with many packaging plastics returning to waste streams in less than a year of manufacture.² However, plastic waste mismanagement continues to be a global challenge, as most plastic waste is either discarded or incinerated. Without action, this issue may compound itself as Geyer *et al.*² project that by 2050 humanity will have produced over 25 000 million metric tons of plastic waste.

1.2. Perspectives of the chemical industry transition

Research regarding a transition of the chemical industry to net-zero emissions has been increasingly gaining attention. Chung *et al.*²⁹ reviewed decarbonisation options for the chemical industry, finding CCU and biomass as significant crosscutting options for the defossilisation of feedstocks as a means to couple the chemical industry with the larger energyindustry system. Material Economics³⁰ develops net-zero



Fig. 4 Conventional naphtha catalytic reforming (CRR) process. The main products from this process are the BTX aromatics, and the by-products include pentane, C_{6+} alkane, liquefied petroleum gas (LPG), and C_{9+} chemicals, with an optional toluene conversion route to increase benzene and xylene yields. Adapted from Jiang *et al.*²⁶

emission pathways for Europe by 2050 for plastics and ammonia, finding for plastics that circular economy, recycling, and bioplastic production are key technologies to reach net-zero emissions without CCS, though they do not consider e-plastic routes with e-methanol as the platform chemical. For ammonia, new processes, *i.e.*, electricity-based ammonia (e-ammonia), are essential for emissions reduction without CCS, as circular economy options are limited. Schneider and Saurat³¹ similarly developed a zero-emission pathway for the European plastic sector by 2050, finding significant roles of plastic recycling, MTO, MTA, and e-naphtha steam cracking to reach net-zero emissions. On a global scale, Kätelhön et al.32 investigated the potential for electricity-based methanol (e-methanol) and electricity-based hydrogen (e-hydrogen) to substitute fossil feedstocks by 2030, emphasising the role of defossilisation of global electricity supply to achieve net-zero emissions in the CCU-based chemical industry. The global net-zero emission chemical industry scenario developed by Saygin and Gielen⁴ still has fossil feedstocks around 25 000 PJ (6940 TWh) in 2050, largely for the production of HVCs, and therefore requires 0.94 GtCO₂/a of fossil CCS and 0.55 GtCO₂/a of bioenergy with CCS (BECCS). Cost-optimal pathways for the global chemical industry without CCS were developed by Zibunas et al.,33



Fig. 5 Deep catalytic cracking of refinery oil for propylene production from wax oil using a zeolite (ZSM-5) catalyst. In addition to the propylene and BTX products, C_4 chemicals, heavy oil, and methane are also produced. Adapted from Zhao *et al.*²⁴ and Gholami *et al.*²⁷

finding that the total energy consumption of the chemical industry ranges from 134 to 160 EJ (37 200–44 400 TWh), and that all primary energies for the global chemical industry can be provided from biomass, ranging from 0 to 65%, and electricity, ranging from 15 to 75% of all resource consumption. Huo *et al.*³⁴ studied the net-zero transition of the global chemical industry, finding a global CO₂ demand of 2.2–3.1 GtCO₂, and a potential supply of 5.2–13.9 GtCO₂ from the power, cement, steel, and pulp and paper sectors; however, this still assumes some fossil usage in power plants and steelmaking, which may not be available in a full defossilisation of the energy-industry system.

While there have been several studies demonstrating the feasibility of the chemical industry to achieve net-zero emissions

at both regional and global scales, there are no studies, to the knowledge of the authors, providing long-term projections of chemical production to the end of the century considering high levels of sustainability. Net-zero transition scenarios for the global chemical industry by 2050 have been performed at a global level,33 and considering fossil CO_2 supply from the power sector and the steel industry at a major region level;³⁴ however, none have been performed at a high regional resolution providing technoeconomic implications for the chemical industry transition in the context of a larger energy-industry defossilisation. Even the leading Integrated Assessment Model (IAM) scenarios, such as that developed by Luderer et al.,³⁵ consider the continued usage of fossil fuels in the chemical industry despite reaching 97.8% RE share in electricity generation. The novelty of this research is therefore in developing the first global chemical and feedstock projections at a high geographical resolution to 2100 for businessas-usual and net-zero emission scenarios considering high and low biomass variations. Furthermore, this research is the first of its kind to evaluate the annualised and levelised costs of the global chemical industry based on green e-ammonia and green e-methanol by applying the levelised cost of electricity (LCOE) according to Bogdanov et al.36 and greenhouse gas (GHG) emission costs according to the 2021 IEA World Energy Outlook.³⁷ This research on a high geographical resolution thus demonstrates specific regional pathways for sustainable feedstocks to enter the global chemical production landscape.

This study is organised as follows: Section 2 describes the methodology and data, Section 3 presents the results, Section 4 provides a discussion of the results, limitations of this study, and recommendations for future work, and Section 5 offers conclusions.

2. Methods and data

This section presents the data and methodology applied for developing scenarios for a defossilised chemical industry. Section 2.1 describes the data and methods applied for developing the chemical production model for the base year of 2020. Section 2.2 explains how future chemical demands were determined and distributed among 145 LUT regions. Section 2.3



Fig. 6 Plastic collection and recycling rates by OECD regions in 2019.²⁸



Table 1Main uses of plastics, shares of plastics by use, and mean productlifetimes according to Geyer et $al.^2$

Plastic use type	Share of total	Mean use lifetime [years]
Packaging	45%	0.5
Building and construction	19%	35
Other	13%	5
Consumer and institutional products	12%	3
Transportation	7%	13
Electrical/electronic	4%	8
Industrial machinery	1%	20

then presents the scenarios that were considered in this research to achieve net-zero CO_2 emissions in the chemical industry.

2.1. Global chemical production landscape

Globally, around 20 chemicals are responsible for 75% of the chemical industry's GHG emissions.⁵ As a starting point, a bottom-up model was developed based on production data for the most downstream chemicals modelled according to Horton,¹⁰ which were then distributed regionally according to Keiner *et al.*³⁸ The bottom-up chemical model developed considers the composition of each primary chemical based on its downstream derivatives according to eqn (2.1), using 2018 production quantities and shares from Horton.¹⁰ Shares of downstream chemical demands for each primary chemical are shown in Table S1 (ESI[†]).

$$P_C = \frac{\left(\sum_i D_{C,i} R_{C,i}\right)}{S_{D,C}} \tag{2.1}$$

where P_C is the total primary chemical demand, $D_{C,i}$ is the downstream chemical demand for chemical *i*, $R_{C,i}$ is the primary chemical demand to produce one ton of chemical *i*, and $S_{D,C}$ is the total share of downstream chemicals modelled relative to the total primary chemical demand.¹⁰

The final chemical demands were separated by their end use of plastics, non-plastics, and pesticides. Plastics as a share of each final chemical were determined according to shares reported by Levi and Cullen and are shown in Table S2 (ESI[†]). The shares of plastics for each final chemical were then used as a basis for determining available plastics for recycling and waste incineration. The amount of plastic chemicals was then determined according to eqn (2.2). Regional plastic collection and recycling rates were applied according to the OECD data and are shown by OECD regions for each NZE scenario in Tables S4–S7 (ESI†), with Table S3 (ESI†) showing the historical CAGR of recycling collection rates by major regions from 1990 to 2019. Similarly, regional plastic incineration growth rates are shown in Table S8 (ESI†), and regional growth rates by scenarios are shown in Tables S9–S12 (ESI†). Collection-to-recycling rates were varied linearly from their regional values in 2019 to reach 100% by the net-zero year for the NZE scenarios, and by the CAGR from 2000 to 2019 under business-as-usual conditions.

$$C_{\text{plastic}} = \sum_{i} C_{\text{final},i} S_{\text{plastic},i}$$
(2.2)

where C_{plastic} is the total chemical-to-plastic demand, $C_{\text{final},i}$ is the total demand of the final chemical *i*, and $S_{\text{plastic},i}$ is the share of the final chemical *i* to plastic production.

These downstream chemicals, along with chlorine, were then allocated categories based on their chemical structure and end-use, primarily to identify which chemicals are used as plastics. The mass balances of the ecoinvent 3.0 database³⁹ were used to determine the primary chemical demand of the downstream chemicals modelled, which then require primary energy feedstocks. Additionally, secondary plastic production was modelled as 6.3% of the total thermoplastics considered in this study. Considering that feedstocks for HVC production varies significantly by region, feedstocks for HVC production, primarily those for steam crackers, were distributed by major global region for Europe, Eurasia, Middle East and North Africa (MENA), sub-Saharan Africa (SSA), the Southeast Asian Association for Regional Cooperation (SAARC), Northeast Asia, Southeast Asia, North America, and South America on a 145 LUT region-basis as used in Bogdanov et al.36

The regional distribution of feedstocks for HVC production is shown in Table 2 for ethylene and Table 3 for propylene. Steam crackers additionally have a pygas component that contains BTX aromatics, among other aromatic chemicals, and BTX shares in pygas were applied according to Levi and Cullen,⁸ and are shown in Table S13 (ESI[†]). While significant quantities of

Table 2Feedstock shares for steam crackers by major region according
to Horton.10Steam crackers are used as the primary technology for
ethylene production, with propylene and BTX aromatic co-products. The
distribution of co-products per tonne of ethylene is shown in Table S13
(ESI)

	Ethane	Propane	Butane	Naphtha	Gas oil	Other ^a
Europe	11%	8%	8%	66%	7%	0%
Eurasia	8%	5%	5%	67%	15%	0%
MENA	63%	24%	5%	8%	0%	0%
Sub-Saharan Africa	25%	25%	0%	0%	0%	50%
SAARC	6%	3%	2%	77%	10%	3%
Northeast Asia	6%	3%	2%	77%	10%	3%
Southeast Asia	6%	3%	2%	77%	10%	3%
North America	44%	19%	13%	19%	3%	1%
South America	26%	10%	0%	58%	6%	0%

^{*a*} The other feedstock assumed to be naphtha.

Table 3 Feedstock shares for propylene production by major region according to Horton.¹⁰ Shares in this table were primarily used to determine the shares of technology applied to the remaining propylene demand after considering propylene co-products from steam crackers. The distribution of co-products per tonne of propylene is shown in Table S14 (ESI)

	Steam crackers	Deep catalytic cracking	Propane dehydrogenation
Europe	26%	25%	7%
Eurasia	26%	25%	7%
MENA	30%	26%	44%
Sub-Saharan Africa	30%	26%	44%
SAARC	44%	25%	31%
Northeast Asia	44%	25%	31%
Southeast Asia	44%	25%	31%
North America	33%	54%	13%
South America	42%	58%	0%

propylene and BTX aromatics are supplied by steam crackers, especially those with heavier feedstocks, this supply is not enough to meet global demand. Therefore, refinery oil and propane are used as feedstocks for propylene production through deep catalytic cracking and propane dehydrogenation, respectively. Additionally, catalytic reforming of naphtha is used to satisfy the remaining BTX aromatic demand, with p-xylene being the chemical most supplied from refinery naphtha.⁸ The result is that there is often a mismatch between supply and demand of BTX aromatics, especially toluene, though this can be somewhat mitigated by converting toluene to benzene and xylenes through toluene disproportionation and toluene hydrodealkylation.^{8,10} For ammonia and methanol production, regional feedstock data were not available; therefore, global feedstock shares for these chemicals were applied according to Levi and Cullen,⁸ as shown in Fig. 8.

The global flow of these chemicals along with chlorine, as shown in Fig. 9, traces the fossil feedstocks to primary and then downstream chemicals that are then used for fertilisers, pesticides, plastics, resins, and fibres. The results of the bottom-up model for 2020 find that the production of primary chemicals required a total energy and feedstock input of 11 029 TWh_{th}. The total fossil feedstock was then compared to the 2020 results provided by the IEA⁴⁰ and DNV⁴¹ to verify the accuracy of the model with the collected data. According to the 2019 IEA energy statistics,⁴⁰ the most recent year of data available, the total nonenergy use was reported at 38.7 PJ, or 10750 TWh_{th}, a 2.6% difference with the calculated number. When compared to 2020 number published in DNV's 2022 energy transition outlook⁴¹ of 40 EJ, or 11111 TWh_{th}, this study's model results find a -0.7% difference, thus verifying the model functionality for the starting year of 2020. Regionally, China has the largest share of the fossil feedstock demand, as well as Saudi Arabia, Korea, and Belgium and the Netherlands (BNL), and several regions in the United States. The regional fossil feedstock demand is shown in Fig. 10.

2.2. Future alternatives

For many energy sectors, such as power, heat, and transport, the discussion on reaching carbon neutrality is often centred around the concept of 'decarbonisation'; however, for the chemical industry, 'decarbonisation' is impossible due to the requirement of carbon feedstocks to produce all large volume organic chemicals.⁴² Therefore, the development of a net-zero emissions or even negative emissions chemical industry must centre around the concept of 'defossilisation', which requires the introduction of new sustainable carbon feedstocks through sustainable biomass and carbon capture and utilisation (CCU).^{4,32,43-45} The most discussed sustainable feedstock for the global chemical industry has been e-methanol and biomass-based methanol (bio-methanol),46 which can either be used directly or as a feedstock for olefins and aromatics through the methanol-to-olefins (MTO) and methanol-toaromatics (MTA) processes. Indeed, methanol has been discussed as a substitute for oil-based feedstocks and fuels as early as the 1980s, though fossil methane was suggested as the major feedstock.47 Both methanol and ammonia can be synthesised through green e-hydrogen and biomass, and e-methanol and bio-methanol as the central feedstocks would lead to a methanol economy⁴⁸ basis for the current petrochemical industry. In this study, electrolysis based organic syntheses⁴⁹⁻⁵³ are not included in this study due to the early stage technology development and low technology readiness levels (TRL 3-5).

2.2.1. Power-to-ammonia. Research regarding sustainable ammonia has largely fallen into three categories focusing on blue hydrogen, green e-hydrogen, and electrochemical ammonia production. The use of blue ammonia for ammonia production would use carbon capture and storage (CCS) to reduce the emissions of conventional ammonia production, though this process would not capture all related CO2 emissions and life-cycle emission factors may be further limited to 60–85%.^{1,54} Green e-ammonia, conversely, proposes the use of green e-hydrogen for the Haber-Bosch synthesis unit, and a temperature of 480 °C at a pressure of 150 bar is applied in this research.¹³ The use of water electrolysis adds a new water demand of 1.6 tH₂O/ tNH₃ for the water electrolyser that is not present in conventional ammonia production, which may cause an additional water stress in regions experiencing water scarcity.¹¹ The green e-ammonia process, as shown in Fig. 11, consists of two primary subsystems,



Fig. 8 Ammonia (left) and methanol (right) production by feedstock in 2013 according to Levi and Cullen.⁸



Fig. 9 Global flow from feedstocks to chemicals in 2020. Downstream chemicals are categorised according to their use for chemicals-to-plastics or by their chemical structure for non-plastic chemicals. Abbreviations: steam cracker (SC), deep catalytic cracking (DCC), catalytic reforming (CR), on-purpose propylene (OPP), toluene hydrodealkylation (TH), toluene disproportionation (TD), polyethylene (PE), vinyl chloride (VC), ethylene oxide (EO), ethylene glycol (EG), styrene (St), polypropylene (PP), acrylonitrile (Acr), propylene oxide (PO), acetone (Ace), cumene (Cu), and terephthalic acid (TA).

which are the gas subsystem supplying nitrogen and hydrogen, and the ammonia synthesis system consisting of the Haber–Bosch reactor.¹⁷ Of the green e-ammonia options, the power-to-ammonia is the most commercially available, at a TRL of 8–9.¹¹ While there is research investigating direct electrochemical synthesis of ammonia from water and nitrogen under low temperature and low pressure conditions, such ammonia synthesis systems are not yet commercially available.¹⁶

2.2.2. Power-to-methanol. Compared to conventional methanol production, which synthesises methanol from carbon monoxide and hydrogen, a power-to-methanol route could convert carbon dioxide and hydrogen to methanol over a Cu/ZnO/Al₂O₃ catalyst, as shown in Fig. 12. The chosen e-methanol synthesis reactor has an

inlet temperature of 210 °C and 76 bar.¹⁹ The entire process has a water requirement of around 27 tH₂O/tMeOH, for water electrolysis; however, this requirement is lower than conventional methanol production, which requires 90 tH₂O/tMeOH.¹⁹ While the overall synthesis route is similar, e-methanol synthesis from carbon dioxide and e-hydrogen is at a lower TRL compared to the conventional route, currently around TRL 7.²⁰ Power-to-methanol has been widely researched as an alternative to conventional production, due to the wide range of applications for methanol to replace fossil fuels in marine and aviation transportation as well as chemical production as envisioned by Olah *et al.*⁴⁸ and Bertau *et al.*⁴⁶ and methanol derivative syntheses envisioned by Banivaheb *et al.*⁵⁶ Furthermore, pilot power-to-methanol plants from Carbon Recycling International



Fig. 10 Total fossil feedstocks for chemicals by region in 2020. Chemical demands that were met with the fossil feedstock supply based on numbers from Horton.¹⁰ The regional structure adopted as used by Bogdanov *et al.*³⁶



Fig. 11 Power-to-ammonia process diagram with nitrogen supplied from an ASU and hydrogen supplied from electrolysis. Adapted from Smith *et al.*¹¹ and Morgan⁵⁵



Fig. 12 Power-to-methanol process diagram with e-hydrogen supplied by water electrolysis and CO_2 being supplied by direct air capture (DAC) or industrial point sources. In this study, CO_2 from DAC is assumed. The catalyst for the methanol synthesis reactor is $CuO/ZnO/Al_2O_3$. Adapted from Adnan and Kibria²⁰ and Bos *et al.*⁶⁴

in Iceland have operated since 2012, and additional pilot plants are being developed by Power to Methanol Antwerp⁵⁷ and Project Air,⁵⁸ which plan to have operational plants by 2022 and 2026, respectively. The first commercial power-to-methanol plant using atmospheric CO₂, with a capacity of 110 ktMeOH, started production in October 2022 in Anyang, Henan Province, China. It uses the emissions-to-liquids technology developed by Carbon Recycling International.⁵⁹

In this research, a carbon dioxide demand of 1.46 tCO_2 / tMeOH is considered to be supplied by a direct air capture (DAC) unit;^{19,60,61} however, carbon dioxide can also be supplied from process emissions from the cement mills, pulp and paper mills, or waste incinerators burning biomass or municipal solid waste.⁶² Techno-economic assessments of the power-tomethanol route have been investigated for a range of carbon inputs including CO₂ from a biogas treatment plant and a fossil ammonia plant,⁶³ DAC,^{60,64} carbon recycling,^{65,66} and other point sources.⁶⁷⁻⁶⁹ Research has also investigated sourcing carbon dioxide from lignite power plants;⁷⁰ however, this would not be a fully sustainable solution given the use of coal as an input for electricity generation and leakage emissions, as point source carbon capture from coal power plants is typically designed around a 90% CO₂ efficiency.⁷¹ Additionally, the fossil carbon embedded in the methanol could return to air or water as CO₂ or other GHG emissions at the end of its life cycle.

2.2.3. Power-to-methanol-to-olefins. Although there are pathways to directly synthesise ammonia and methanol from green e-hydrogen, such routes are not readily available for olefins or aromatics, as the single stage conversion of hydrogen and CO₂ to olefins is still at a TRL of 3-4.43 Therefore, the conversion of methanol has been proposed to substitute oil feedstocks. The MTO process, however, has largely been investigated to use coal-based methanol as an input, largely in China due to the high availability of coal.^{24,72} Due to its commercialisation in China, MTO already has a high TRL of 8-9.43 MTO, as shown in Fig. 13, operates at around 500 °C and 2.5 bar over a SAPO-34 type catalyst, with a carbon selectivity ranging from 78 to 82%.²⁴ The methanol input for MTO assumed in this study is 16.34 MWh_{MeOH,LHV}/tOlefin.⁶ Multiple MTO processes have been developed, and different ratios of ethylene and propylene can be achieved depending on the catalyst used.73 In addition to the ethylene and propylene products, there is a heat by-product of 0.688 MWh_{th}/tOlefin at 500 °C and a water by-product of 1.685 tH₂O/tOlefin.⁷⁴

2.2.4. Power-to-methanol-to-aromatics. Compared to the MTO route, MTA is much less developed, with a TRL of 7.⁴³ MTA has also largely been researched in the context of coalbased methanol.^{26,76,77} The MTA process, as shown in Fig. 14, converts methanol to aromatics over a zeolite catalyst, HZSM-5, at 370–540 °C and 20–25 bar.⁴³ Of the final products, BTX



Fig. 13 Power-to-methanol-to-olefins process diagram. Methanol synthesis occurs over a CuOZnO/Al₂O₃ catalyst and the MTO reactor uses the SAPO-34 catalyst. Adapted from Jiang *et al.*²⁵ and Dimian and Baldea.⁷⁵



Fig. 14 Power-to-methanol-to-aromatic process diagram. Methanol synthesis occurs over a CuOZnO/Al₂O₃ catalyst, and methanol aromatisation occurs over a HZSM-5 catalyst. Adapted from Jiang *et al.*²⁶

aromatics compose around 16% of the total yield by weight; therefore, a significant methanol input of 34.46 MWh_{MeOH,LHV} is required.⁷⁶ The most significant by-products of the MTA process are liquefied petroleum gas (LPG), which is produced at a rate of 1.24 tLPG/tBTX, and water, which is produced at a rate of 3.224 tH₂O/tBTX.⁷⁶ Additionally, the MTA process is highly exothermic, with a heat output of 2.838 MWh_{th}/tBTX.⁷⁴

2.2.5. Fossil feedstocks with carbon capture and storage and biomass feedstocks. Considering the composition of emissions, the IEA reports that the highest share of emissions from primary chemical production comes from ammonia, at 49%, followed by HVCs, at 27%, and methanol, at 24%.³ For methanol and ammonia, a high share of emissions comes from the

reduction of the fossil feedstock to syngas. Therefore, CCS has been proposed to remove the high amount of process CO_2 emissions, and, for ammonia production, some carbon capture is already utilised for downstream urea production.^{1,78} Conversely, due to the conversion of fossil hydrocarbons to HVCs, most emissions are a result of process energy, as feedstock losses, and thus feedstock emissions, tend only to be 0.5% of the feedstock input.²² For these HVC production routes, CCS would largely be unnecessary if the process energy inputs were decarbonised. Gabrielli *et al.*⁴² highlighted the potential for CCS to reduce emissions in the chemical industry; however, due to the point source capture efficiency of 90%, additional DACCS is required to capture the remaining 10% of CO₂ that is

not captured in the CCS stage. Furthermore, while fossil carbon may not be emitted to the atmosphere during the chemical production stage, it may be emitted at the end-of-life stage through waste incineration or degradation in landfills. Additionally, fossil chemicals with CCS may lead to a fossil lock-in for the chemical industry, and lead to stranded fossil assets as fossil reserves become increasingly nonviable⁴² as well as increased life-cycle emissions from the carbon content of fossil-based chemicals. Therefore, to avoid potential fossil lock-in and related non-CO₂ air pollutants,⁷⁹ fossil-CCS options are excluded from this analysis.

Biochemical routes have also been widely suggested to substitute fossil feedstocks for ammonia,⁸⁰⁻⁸² methanol,^{65,83-85} olefins,^{24,86-88} and aromatics,⁸⁹⁻⁹¹ as part of a larger bioeconomy.⁹² While biochemical routes are available, increased scrutiny must be placed on the sourcing of the biomass feedstocks. Today, bioethylene is produced in Brazil via bioethanol dehydration using sugarcane as the biomass feedstock for ethanol production.⁹³ However, as biomass resources become increasingly limited, strict sustainability requirements must be placed on biomass resource use, as biomass competes both with food and feed supply and bioenergy in other energy sectors such as biofuels for transport. Globally, the sustainable bioenergy limit has been estimated to be 100 EJ/a⁹⁴ (27 800 TWh). Sustainable biomass use effectively eliminates first-generation biomass from being used for chemical production, and, therefore, only biomass residues and wastes, e.g., lignocellulosic biomass, should be considered for biochemical production.

2.3. Future primary chemical production and plastic waste management

To determine future chemical demands, individual CAGRs were attributed to each downstream chemical, derived from the IEA⁵ and Kätelhön *et al.*³² These CAGRs were then scaled according to the gross domestic product (GDP) per capita chemical demand model, based on the LUT-EEES macro-economic scenario

of Keiner *et al.*³⁸ from 2020 to 2100 in 10 years intervals and distributed among the 145 LUT regions. Using the shares of chemicals-to-plastics as established in Section 2.1, historical plastic production data from the OECD,²⁸ and plastic lifetime distributions from Geyer *et al.*² (Table 1), the available plastic waste in a given year was determined. Plastics were then separated by collection for recycling and waste incineration. The collection-to-recycling rate was linearly varied from today's levels by major region to 100% by the net-zero emission year. Mechanical recycling was assumed to substitute primary plastic chemicals with secondary plastics at a carbon efficiency of 98% and electricity consumption of 0.234 MWh_{el}/t_{plastic,out}.⁹⁵ Fossil chemicals-to-plastics that are incinerated result in positive GHG emissions, corresponding to the carbon content of chemicals used for plastic production.³⁹

The total plastic production and plastics by application are shown in Fig. 15. For all non-plastic chemicals, only primary production methods were applied, due to the lack of relevant recycling options. Additionally, the carbon content of nonplastic fossil chemicals is included in the GHG emissions reporting in their respective production years as CO_{2ea} emissions. In the net-zero emissions chemical industry scenarios defined in Section 2.5, the maximum plastic recycling rates were set to 60% in the net-zero emission year, evolving from today's regional recycling rates, as shown in Fig. 6. While plastic recycling rates could in theory be higher, especially with chemical, or back-to-monomer, recycling, this recycling target is in line with similar research, e.g., Saygin and Gielen,⁴ who establish a 66% plastic recycling target by 2050. Chemical recycling, which can reduce plastics either to monomers, feedstock oils, or even CO/H₂ syngas, may provide alternative recycling routes for plastics that might be challenging to recycle mechanically,⁹⁶ though it is not applied in this analysis. The physical parameters for each chemical production route are presented in Table S19 (ESI[†]), due to the high number of processes considered in this study.



Fig. 15 Total plastic production (line) and plastics by applications (bars) from 1990 to 2100 in 10-year intervals.

2.4. Economics of chemical production

Annualised costs and levelised costs of primary chemicals (LCOC) for the scenarios defined in Section 2.3 were then determined to analyse the economic viability of green chemical pathways compared to business-as-usual conditions. A weighted average cost of capital (WACC) of 7% was applied for all technologies, and the capital recovery factor (crf) was determined using eqn (2.3). These costs were determined according to eqn (2.4) for annualised costs and eqn (2.5) for LCOC.

$$\operatorname{crf}_{i} = \frac{\operatorname{WACC}(1 + \operatorname{WACC})^{n_{i}}}{(1 + \operatorname{WACC})^{n_{i}} - 1}$$
(2.3)

$$Cost_{Annualised} = \sum_{i} (Capex_{i} \cdot crf_{i} + Opex_{fix,i} + Opex_{var,i}) + Cost_{fuel} + Cost_{El} + Cost_{GHG}$$
(2.4)

$$LCOC = \frac{Cost_{Annualised}}{C_{primary} + C_{secondary}}$$
(2.5)

where crf is the capital recovery factor for technology *i*, WACC is the weighted average cost of capital, *n* is the lifetime of technology *i*, Capex_i is the capital expenditures for technology *i*, Opex_{fix,i} is the fixed operational expenditures for technology *i*, Opex_{var,i} is the variable operational expenditures for technology *i*, Cost_{fuel} is the total fuel costs, Cost_{El} is the total electricity cost, Cost_{GHG} is the total GHG emissions cost, C_{primary} is the total production of primary chemicals, and $C_{\text{secondary}}$ is the total secondary chemical production from plastic recycling.

Furthermore, costs were determined according to major regions to identify which regions may have the best conditions for low-cost green chemicals. Fossil fuel feedstock prices were set according to Bogdanov *et al.*³⁶ Two levelised costs of

electricity (LCOE) were applied, one for 'direct' electricity demands required to operate DAC and chemical production plants that were assumed to operate at 8000 h/a, and another for 'indirect' electricity used by water electrolysers to produce e-hydrogen, which can operate flexibly according to solar PV and wind generation profiles. In major regions, direct LCOE was taken from Bogdanov et al.,36 as embedded in the energyindustry system, and indirect LCOE was determined by considering a hybrid solar PV-wind power plant using utility-scale shares of solar PV and wind power from Bogdanov et al.36 from 2020 to 2050 in 10-year intervals. Additional hydrogen storage was then required assuming a storage requirement of 500 GWh_{H2,LHV} per baseload GWh_{H2,LHV} with an energy-topower ratio of 200.97 GHG emissions costs were varied from 50, 114, 180, and 220 €/tCO₂, as set in the 2021 IEA World Energy Outlook.³⁷ Biomass costs were set to 34.7 € per MWh_{th} assuming a wood pellet price of 200 USD/t⁹⁸ and an exchange rate of 1.2 USD per €. A full list of financial assumptions is listed in Tables S11-S13 (ESI[†]).

2.5. Scenarios for defossilised chemicals

Seven scenarios were then established to study rapid and delayed defossilisation pathways compared to business-asusual (BAU) conditions considering high and low biomass variations, as well as a transition to direct electric process heat, which are defined in Table 4. In the BAU scenario, plastic recycling rates increased according the CAGR of plastic recycling from 2010 to 2019,²⁸ whereas in the net-zero emission (NZE) pathways these were varied to reach a maximum recycling rate of 60% in their net-zero emission year. Similarly, plastic incineration rates were varied according to historical CAGR rates²⁸ in the BAU scenario, and linearly in the NZE

 Table 4
 Technology transition assumptions for each scenario studied from fossil routes (conventional) to those using renewable electricity and sustainable biomass resources (improved)

		2020	2030	2040	2050	2060	2070	2080	2090	2100
BAU	Conventional	100%	100%	100%	100%	100%	100%	100%	100%	100%
	Improved	0	0	0	0	0	0	0	0	0
	Biochemical share	0	0	0	0	0	0	0	0	0
NZE2050	Conventional	100%	85%	40%	0	0	0	0	0	0
	Improved	0	15%	60%	100%	100%	100%	100%	100%	100%
	Bioplastic share high	0	10%	25%	40%	50%	55%	60%	60%	60%
	Bio-ammonia/bio-methanol share high	0	5%	10%	15%	20%	20%	20%	20%	20%
	Bioplastic share low	0	2%	5%	10%	10%	10%	10%	10%	10%
	Bio-ammonia/bio-methanol share low	0	1%	3%	5%	5%	5%	5%	5%	5%
	Direct electricity heating substitution	0	11.9%	88.1%	99.8%	100%	100%	100%	100%	100%
NZE2040	Conventional	100%	50%	0	0	0	0	0	0	0
	Improved	0	50%	100%	100%	100%	100%	100%	100%	100%
	Bioplastic share high	0	15%	30%	50%	55%	60%	60%	60%	60%
	Bio-ammonia/bio-methanol share high	0	10%	15%	20%	20%	20%	20%	20%	20%
	Bioplastic share low	0	5%	10%	10%	10%	10%	10%	10%	10%
	Bio-ammonia/bio-methanol share low	0	3%	5%	5%	5%	5%	5%	5%	5%
	Direct electricity heating substitution	0	50.0%	98.2%	100%	100%	100%	100%	100%	100%
NZE2060	Conventional	100%	90%	66%	25%	0	0	0	0	0
	Improved	0	10%	34%	75%	100%	100%	100%	100%	100%
	Bioplastic share high	0	5%	15%	25%	35%	45%	55%	60%	60%
	Bio-ammonia/bio-methanol share high	0	5%	8%	10%	15%	20%	20%	20%	20%
	Bioplastic share low	0	1%	2%	4%	8%	10%	10%	10%	10%
	Bio-ammonia/bio-methanol share low	0	1%	2%	3%	4%	5%	5%	5%	5%
	Direct electricity heating substitution	0	0.2%	11.9%	88.1%	100%	100%	100%	100%	100%

scenarios to reach 40% by the net-zero emission year. Therefore, the NZE scenarios establish plastic waste management leading to no landfilling nor mismanagement of plastic waste. The transition is modelled to transition from conventional fossil routes (Fig. 10) to one based on renewable methanol and ammonia.

The main variation among NZE scenarios, thus, is the primary energy input to these primary platform chemicals, which can be either renewable electricity (e-chemicals) or biomass (bio-chemicals). Therefore, each scenario was varied among high (H) and low (L) biomass scenarios, e.g., NZE2050H. For bio-chemical routes, no first-generation energy crops were considered as feedstocks, due to potential land-use constraints and conflicts with food production. Consequently, only lignocellulosic second-generation biomass was considered for biochemical production, with a maximum sustainable biomass of 100 EJ⁹⁴ (27 800 TWh). In the high biomass scenarios, the bioenergy limit was set to 70 EJ (19400 TWh), and in the low biomass scenarios, the bioenergy limit was set to 25 EJ (6440 TWh). Bio-chemicals were prioritised for plastics, with smaller shares for bio-methanol and biomass-based ammonia (bio-ammonia). For non-plastic HVCs, no biomass feedstocks were applied due to the high amounts of biomass needed for bioplastics. The scenarios are defined as follows:

Business-as-usual (BAU). Fossil-based chemical production continues to the end of the century, maintaining today's levels of plastic recycling. Process heating is supplied by natural gas, and power sector emissions are assumed to reach zero by 2050, assuming the defossilisation of the power sector, but not of the chemical industry.

NZE 2050H/NZE 2050L. Feedstocks to the global chemical industry in this scenario are completely defossilised by 2050, with high (H) and low (L) biomass feedstock variations. Plastic recycling rates reach 60% in 2050 and power sector emissions reach net-zero emissions by 2050.

NZE 2040H/NZE 2040L. Feedstocks to the global chemical industry in this scenario are completely defossilised by 2040, with high (H) and low (L) biomass feedstock variations. Plastic recycling rates reach 60% in 2040 and power sector emissions reach net-zero emissions by 2040.

NZE 2060H/NZE 2060L. Feedstocks to the global chemical industry in this scenario are completely defossilised by 2060, with high (H) and low (L) biomass feedstock variations. Plastic recycling rates reach 60% in 2060 and power sector emissions reach net-zero emissions by 2060.

3. Results

The results of this research are organised as follows: Section 3.1 presents the future global chemical production landscape from 2020 to 2100 in 10 year intervals, Section 3.2 examines the energy system and feedstock requirements for each scenario, Section 3.3 shows the evolution of regional chemical production due to increased GDP/capita in the emerging economies of the world, and Section 3.4 considers the annualised costs and

LCOC of a global chemical transition to e-chemicals and biochemicals.

3.1. Future chemical demand

Under BAU conditions, the total primary volume of the 21 chemicals modelled increases significantly from 1220 Mt (10069 TWh_{LHV}) to 2763 Mt (22092 TWh_{LHV}) in 2050 and 3481 Mt (27795 TWh_{LHV}) in 2100, as shown in Fig. 16.

With increased plastic recycling, however, primary chemical production can be reduced by 8.1%, 8.1%, and 5.9% in 2050 for NZE 2050, NZE 2040, and NZE 2060, respectively. By 2100, primary chemical production will be reduced by 6% compared to BAU across all NZE scenarios. The effect of recycling can be most observed in the demands of ethylene and propylene, due to the high shares of olefins in global plastics, whereas the demands of BTX aromatics are only slightly reduced due to their high use in non-plastic applications. The primary ethylene demand in the BAU scenario, for example, reaches 358 Mt in 2050, which is 12% higher than the ethylene demand in the NZE 2050 scenarios, at 296 Mt in 2050. Increased levels of both plastic recycling and stabilisation of GDP/capita levels across global regions will lead to a levelling off of global chemical demands in the latter half of the century, especially from 2070 to 2100.

This effect can be better observed in Fig. 17, where chemical demands by end use are shown.

In 2050, secondary plastics will be composed of 30.7%, 30.7%, and 25.9% of the total plastic demand in the NZE 2050, NZE 2040, and NZE 2060 scenarios, respectively, and reach 53.7% of the global plastic demand by 2100 across scenarios, compared to 40.8%% in the BAU scenario. The total savings of primary final chemical production in the NZE 2050 scenario amounts to 223 Mt in 2050 and 210 Mt in 2100 compared to the BAU scenario. The rapid increase of secondary plastic shares is largely due to the high shares of packaging plastics, which typically have a lifetime of less than a year and are thus available for recycling within the same annual time step.

3.2. Flows and feedstocks for the future chemical industry

By 2050, the significant increase in global chemical demands implies similar increases in hydrocarbon feedstocks, which can either result in significant increases in fossil feedstocks, as shown in Fig. 18, or a transition to renewable hydrocarbon feedstocks, as shown in Fig. 19 and Fig. 20. In the BAU scenario, the total fossil feedstock demands increase from 11 029 TWh_{th,LHV} to 23 591 TWh_{th,LHV} in 2050, corresponding to a CAGR of 2.6%. The steam cracker remains the key technology to produce HVCs, consuming 54.1% of the total fossil feedstocks. Under BAU conditions, the highest shares of losses come from conventional ammonia and methanol production, whereas HVC production mostly results in byproducts, whose energy content reaches 7320 TWh_{th,LHV}, with only 0.5% feedstock losses.²²

Comparatively, the NZE 2050H and NZE 2050L scenarios present a full transition of primary energy inputs from today's

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fossil fuel structure to ones dominated by renewable electricity and biomass, as in the case of the NZE 2050H scenario. By 2050, 27759–33031 TWh_{el} and 2548–9848 TWh_{th} of biomass feedstocks are required to satisfy the global chemical feedstock demands. In NZE 2050H, the biomass demand corresponds to 35.4% of the global sustainable biomass demand, and, in NZE 2050L, only 9.2% is used. Methanol becomes the most important chemical feedstock as all olefins and BTX aromatics are derived from e-methanol and bio-methanol. While the MTO process shows a high energy conversion from the methanol



Fig. 18 Global chemical flows in 2050 for the BAU scenario.



feedstock to olefin, the MTA process has significant shares of input energy content converted to by-products, most notably LPG. Total losses in NZE 2050L are lower than those of NZE 2050H despite significantly increased electrolyser losses, largely due to the higher energy efficiency of the power-to-methanol process compared to biomass-to-methanol route. The new dominant power-to-chemical structure additionally introduces a new excess heat by-product, which can either be used in district heating systems, CO_2 DAC, or used for other industrial process heating demands as the excess heat of the MTO process is at 500 °C. Additionally, hydrogen by-products from chlorine production and the MTO process in this study are assumed to partially substitute hydrogen from electrolysis, although the total impact of these hydrogen by-products is rather small. The energy content of recycled plastics reaches 1961 TWh_{th} in 2050, and 30.7% of total chemicals-to-plastics. Detailed Sankey diagrams from 2020 to 2050 in 10 year intervals, 2070, and 2100 for BAU, NZE 2050H, and NZE 2050L are shown in Fig. S5–S16 (ESI†).







Examining the feedstock transition pathway in more detail as shown in Fig. 21, the total feedstock demand for chemicals can be expected to increase similarly with the growth in chemical demands; however, increased plastic recycling has significant effects on reducing feedstock demands. In the BAU scenario, fossil feedstock demands increase from 11 029 TWh_{th} in 2020 to 23 591 TWh_{th} in 2050, and finally 27 705 TWh_{th} in 2100. Comparatively, total feedstock demands in the NZE 2050 scenarios reach 20 628 TWh_{th} and 25 515 TWh_{th} in 2100, with varying levels of e-methanol and bio-methanol and ammonia. According to the scenario definition, the largest increases of renewable feedstocks occur from 2030 to 2040 and 2040 to 2050, which will be required if the global chemical industry is to reach net-zero emissions by 2050.

Furthermore, significant e-chemical feedstocks imply significant increases in renewable electricity capacity, as the total electricity demands for the NZE 2050 scenarios reach 35 075-40 169 TWh_{el} in 2050 and 40 593-50 257 TWh_{el} in 2100 for the NZE 2050H and NZE 2050L scenarios, respectively, from just 1645 TWhel in 2020. For reference, the IEA reported that the total electricity generation in 2020 was 26708 TWhel.⁹⁹ Therefore, by 2050, the chemical industry could be expected to have electricity demands that are 131-150% of today's global electricity generation. This result suggests that the massive ramping of renewables, especially solar PV and wind power, will be required for the defossilisation of chemicals, as supplying this electricity demand with today's electricity mix would undoubtedly cause a massive increase in chemical industry emissions.³² The highest quantities of electricity will be needed for e-hydrogen production, reaching 31025 and 41 029 TWhel in the NZE 2050H and NZE 2050L scenarios, respectively, as shown in Fig. 22. Comparatively, direct electricity demands remain low across scenarios, only reaching 2564 (96% of total electricity demand), 6449 (13%), and 5523



Fig. 22 Direct (for chemical production plants), indirect (for water electrolysis), and heating electricity demands for the BAU (top left), NZE 2050H (top right), and NZE 2050L (bottom) scenarios.

(11%) TWh_{el} in 2100 in the BAU, NZE 2050H, and NZE 2050L scenarios, respectively.

A small but important shift occurs with process heating demands, as they transition from fossil boilers to electric boilers and heat pumps, as shown in Fig. 23. In the BAU scenario, continued usage of fossil chemical processes leads to the highest energy requirements for heat of the scenarios studied, increasing from 1417 TWh_{th} in 2020 to 3265 TWh_{th} in 2050, and 3521 TWh_{th} in 2100. Comparatively, the highest share of future heat demands for the chemical industry in the NZE scenarios is the heat required CO₂ via DAC using heat pumps for the powerto-methanol process. The total energy for heating in the NZE 2050 scenarios ranges from 2610 to 2919 TWh in 2050 and 7250 to 9420 TWh in 2100 for NZE 2050H and NZE 2050L, respectively. Other process heating demands for primary and downstream chemical production then transition to direct electric heating, assuming that temperature requirements are higher than what can be supplied by heat pumps for the downstream production processes. By 2100, 1549-1581 TWh_{th} of recoverable heat will be available.

The impact of the transition from fossil to sustainable feedstocks on emissions is shown in Fig. 24 by type for the BAU, NZE 2050H, and NZE 2050L scenarios.

Fig. 25 then shows emissions levels for all scenarios studied. Emissions from the production of the chemicals considered in this study were found to be 1240 MtCO2eq in 2020, corresponding to 3.6% of global CO₂ emissions.⁹⁹ The fossil content of non-plastic chemicals and incineration of fossil plastic waste contribute an additional 685 MtCO_{2eq}, leading to total emissions of 1926 MtCO_{2eq}. In the BAU scenario, emissions steadily increase in the coming decades, reaching 3769 MtCO_{2eq} in 2050, before levelling off at an emissions level of 5713 MtCO_{2eq} in 2100. Interestingly, the large increase of production-related emissions in the BAU scenario seems to largely be a result of emissions from process heat, rather than from feedstocks. If all process emissions were reduced to zero in the BAU case, production-related emissions would only reach 1626 MtCO2eq by 2100. Furthermore, the growing levels of fossil plastic waste and fossil chemical production lead to significant growth in the GHG impact of fossil chemicals.

While the NZE 2050 scenarios achieve emissions targets by the target year, it is not due to a steady decline in emissions. Rather, an emissions peak in 2030 due to increased fossil feedstock usage and rising electricity demands in an electricity system environment with an emissions factor of $0.086 \text{ tCO}_2/\text{MWh}_{el}$, with the NZE 2040 scenarios being the



Fig. 23 Process heating demands and excess heat from exothermic reactions in the power-to-methanol, MTO, and MTA processes from 2020 to 2100 for BAU (top left), NZE 2050H (top right), and NZE 2050L (bottom) scenarios. Note that the value listed for heat pumps is the electricity input and is thus in TWh_{el}.

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Fig. 24 Chemical industry emissions by source for the BAU (top left), NZE 2050H (top right), and NZE 2050L (bottom) scenarios. In the NZE 2050 scenarios, low levels of GHG emissions are still present from the incineration of fossil plastic waste in 2050–2080, despite net-zero emission chemical production.



exception. After the first time-step, emissions decline from 2030 to 2040 before reaching net-zero CO_2 emissions in 2050. Comparing the emissions trajectories of all scenarios shows that the high biomass scenarios tend to have lower emissions in the transition years relative to their low biomass counterparts. This is again likely due to the increased electricity usage of the low biomass scenarios.

3.3. Geographical distribution

While global feedstock and process energy demands are projected to increase, regional pathways were found to vary substantially. However, all global regions experience massive growth in electricity and hydrogen demands, with the regions of the Global South having a much larger role in global chemical production compared to today's regional structure (Fig. 10). The regional structures of key inputs to the global chemical industry for the BAU and NZE 2050 scenarios are shown in Fig. 26. The regional results show that the increases in total demand for electricity, fossil feedstocks in the BAU scenario, and e-hydrogen in the NZE 2050 scenarios is primarily driven by significant growth in the chemical demand in Africa, South America, and Southeast Asia. The NZE 2050H scenario, despite high biomass usage for plastics, finds regional electricity demands above 200 TWh_{el} in 2050, and, correspondingly, the highest chemical demand regions experience e-hydrogen demands up to 930 TWh_{H2,LHV}. In the NZE 2050L scenario, these demands are even higher on a regional basis, as the average regional electricity and hydrogen demands are 277 TWh_{el} and 159 TWh_{H2,LHV}, with Central China having the highest of both demands at 1919 TWh_{el} and 1104 TWh_{H2,LHV}, respectively.

While industrialised regions of the world including Europe and North America experience only limited growth in the total feedstock required, as shown in Table 5, e-hydrogen and biomass demands substantially grow across all major regions. Table 6 and Table 7 indicate that for many major regions, peak e-hydrogen demands for chemicals will occur around 2050 and 2060, though the regions of Eurasia, MENA, sub-Saharan Africa, and North America experience peak e-hydrogen demands in 2100. Due to increasing shares of biomass feedstocks in the NZE 2050 scenarios, many regions do not have their peak

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Fig. 26 Regional total electricity and fossil feedstocks for the BAU scenario (top left), and e-hydrogen demands (middle and bottom right) for the BAU (top), NZE 2050H (middle), and NZE 2050L (bottom) scenarios.

biomass demands until 2100. Peak process electricity demands largely follow the trends of e-hydrogen demands, largely due to the regional co-location of MTO and MTA plants and electrolyser capacities. Geographical results thus further highlight a shift of chemical production from the Global North to the Global South as the sub-Saharan Africa and SAARC regions experience the most rapid growth in renewable feedstocks and will have the highest e-hydrogen and biomass feedstock demands by 2100. Additional regional results by scenario for 2030, 2040, 2070, and 2100 are available in Fig. S17–S49 (ESI†).

3.4. Cost structure for a green chemical transition

The annualised cost structures of the BAU, NZE 2050H, and NZE 2050L scenarios by cost component and by major region are shown in Fig. 27. Due to the lack of available cost data for downstream processes, the annualised costs were developed for

primary production of the main platform chemicals and chlorine, as well as secondary plastic production, to represent the costs of feedstock substitution. The total annualised cost of the chemical industry in 2020 was found to be 630 b€, with fuel and feedstock costs alone totalling to 369 b€. Under BAU conditions, this cost structure continues, with GHG emission pricing mechanisms adding an additional 805 b€ by 2050 from the continued fossil fuel usage as feedstocks and for process heat. Annualised costs in the BAU scenario will reach 2274 b€ in 2050.

The results suggest that the net-zero emission chemical industry can have lower annualised costs in 2050 compared to those under BAU conditions, when low-cost renewable electricity is widely applied for e-methanol and e-ammonia, as is the case in the NZE 2050L scenario, with annualised costs of 1938 b€. Conversely, the NZE 2050H scenario finds higher

 Table 5
 Projections of primary energy and direct electricity demands for the BAU scenario from 2020 to 2100 in 10-year intervals by major regions. The peak feedstock and electricity demands for each major region are highlighted in bold. Abbreviations: Middle East and North Africa (MENA) and South Asian Association for Regional Cooperation (SAARC)

Fossil feedstock demand BAU [TWh _{th}]	2020	2030	2040	2050	2060	2070	2080	2090	2100
rossn recustock demand DAO [1 Wh _{th}]	2020	2030	2040	2030	2000	2070	2000	2090	2100
Europe	1577	2046	1971	1861	1713	1621	1578	1568	1582
Eurasia	1081	1123	916	770	773	761	746	720	730
MENA	1056	1400	1441	1554	1793	1905	1924	1874	1899
Sub-Saharan Africa	116	672	1906	4180	6577	8230	9086	9441	1006
SAARC	760	2072	3703	5407	6137	6110	5788	5470	5274
Northeast Asia	3205	4717	4818	4379	3828	3488	3286	3149	3082
Southeast Asia	749	1360	1853	2261	2352	2243	2167	2114	2092
North America	2125	2415	2035	1713	1672	1659	1679	1679	1734
South America	362	785	1148	1466	1518	1428	1348	1284	1248
Total global	11 029	16 590	19790	23 591	26364	27446	27 601	27 300	27 70
Direct electricity demand BAU [TWh _{el}]	2020	2030	2040	2050	2060	2070	2080	2090	210
Europe	73	96	112	128	137	140	139	138	136
Eurasia	50	51	50	49	56	59	61	61.1	61.
MENA	56	71	86	109	144	164	174	179	182
Sub-Saharan Africa	6	32	107	274	489	658	768	849	91
SAARC	35	96	204	351	466	513	518	507	49
Northeast Asia	150	221	274	302	320	316	304	291	27
Southeast Asia	35	63	101	146	178	191	193	192	18
	104	116	115	114	129	138	144	147	149
North America	104	110	115	114	145				
North America South America	104 17	36	63	94	115	123	123	120	11

 Table 6
 Projections of renewable feedstocks and direct electricity demands for the NZE 2050H scenario. The peak feedstock and electricity demand for each major region are highlighted in bold

- Total e-Hydrogen demand NZE 2050H [TWh _{H2,}	_{LHV}] 20	20 2030	0 2040	2050	2060	2070	2080	2090	2100
Europe	0	220	1066	1556	1339	1256	1217	1229	1256
Eurasia	0	119	451	560	541	545	560	574	591
MENA	0	121	767	1308	1410	1468	1489	1528	1576
Sub-Saharan Africa	0	65	1026	3480	5038	6079	6556	7102	7713
SAARC	0	214	1976	4471	4754	4623	4281	4106	4025
Northeast Asia	0	495	2607	3748	3137	2810	2625	2568	2561
Southeast Asia	0	141	974	1828	1780	1706	1619	1601	1609
North America	0	261	1051	1288	1212	1262	1312	1361	1416
South America	0	87	610	1191	1152	1095	1017	984	970
Total global	0	1724	4 1052	8 1943	1 20362	20 844	20 675	21054	21718
Total Biomass demand NZE 2050H [TWh _{th}]	2020	2030	2040	2050	2060	2070	2080	2090	2100
Europe	0	31	284	729	788	796	845	852	873
Eurasia	0	17	113	230	291	329	385	395	406
MENA	0	22	209	638	920	1060	1184	1191	1211
Sub-Saharan Africa	0	11	309	1935	3757	5026	5901	6197	6577
SAARC	0	31	576	2413	3389	3579	3531	3255	3118
Northeast Asia	0	71	708	1796	1874	1768	1781	1724	1716
Southeast Asia	0	20	276	947	1205	1251	1274	1229	1219
North America	0	37	268	534	646	772	917	955	995
South America	0	12	173	627	791	812	805	755	731
Total global	0	251	2916	9848	13 661	15 392	16 622	16554	16 846
Direct Electricity demand NZE 2050H [TWh _{el}]	2020	2030	2040	2050	2060	2070	2080	2090	2100
Europe	73	135	259	367	355	347	347	346	349
Eurasia	50	72	109	128	139	147	156	159	162
MENA	56	99	192	313	387	426	450	456	464
Sub-Saharan Africa	6	45	258	872	1457	1863	2101	2248	240
SAARC	35	135	491	1108	1354	1390	1336	1261	121
Northeast Asia	148	308	636	889	837	778	745	717	703
Southeast Asia	35	88	241	446	497	502	493	481	475
North America	104	161	254	294	312	341	369	380	392
				202	222	323	312	297	287
South America	17	51	150	292	323	323	312	297	207

 Table 7
 Projections of renewable feedstocks and direct electricity demands for the NZE 2050L scenario. The peak feedstock and electricity demand for each major region are highlighted in bold

Total e-Hydrogen demand NZE 2050L [TWh _{H2,}	_{LHV}] 2020	2030	2040	2050	2060	2070	2080	2090	2100
Europe	0	233	1179	1826	1653	1580	1567	1582	1618
Eurasia	0	126	496	646	656	680	719	738	760
MENA	0	130	850	1547	1779	1902	1982	2023	2079
Sub-Saharan Africa	0	70	1150	4211	6554	8142	9015	9682	10 45
SAARC	0	227	2206	5380	6117	6089	5750	5458	5320
Northeast Asia	0	523	2888	4417	3883	3529	3362	3283	3273
Southeast Asia	0	149	1085	2183	2263	2218	2149	2112	2116
North America	0	276	1157	1486	1469	1577	1693	1758	1829
South America	0	92	679	1426	1470	1427	1351	1298	1274
Total global	0	1825	11690	23 122	25844	27 143	27 589	27 935	28 720
Total Biomass demand NZE 2050L [TWh _{th}]	2020	2030	2040	2050	2060	2070	2080	2090	2100
Europe	0	6	60	188	162	151	149	150	153
Eurasia	0	3	24	60	60	63	68	69	71
MENA	0	4	44	166	189	201	207	208	212
Sub-Saharan Africa	0	2	65	501	773	947	1027	1080	114
SAARC	0	6	122	625	697	676	619	571	547
Northeast Asia	0	14	150	463	385	336	314	304	302
Southeast Asia	0	4	58	245	248	237	224	216	214
North America	0	7	57	138	133	147	161	167	174
South America	0	2	37	162	163	154	141	133	128
Total global	0	50	620	2548	2811	2910	2909	2899	2948
Direct Electricity demand NZE 2050L [TWh _{el}]	2020	2030	2040	2050	2060	2070	2080	2090	2100
Europe	73	134	245	331	314	305	301	300	301
Eurasia	50	71	104	116	124	129	135	137	139
MENA	56	97	181	281	339	369	385	391	397
Sub-Saharan Africa	6	45	242	775	1256	1590	1776	1907	2043
SAARC	35	133	460	988	1174	1196	1142	1083	104
Northeast Asia	149	307	601	800	739	683	648	623	609
Southeast Asia	35	87	226	399	434	434	423	413	408
North America	104	160	241	268	278	300	319	328	337
South America	17	50	141	261	282	280	267	255	247
Total global	524	1086	2447	4709	5663	6118	6309	6347	6449

annualised costs of 2147 b€, which is primarily due to the high biomass feedstock costs, which total 342 b€ in 2050. Due to the widespread application of power-to-chemicals routes, the annualised cost structure transitions from being fuel dominated to being driven by electricity costs and capital expenditures (capex), especially for water electrolysers and hydrogen storage, as shown in Fig. S31 (ESI†). Increases in operational expenditures (opex) also experience a noticeable increase, largely due to high opex_{fix} for hydrogen storage and chloralkali electrolysers. Furthermore, increased plastic recycling plays a limited but important role in reducing costs for primary chemical feedstocks.

Regional contributions to global annualised costs follow the structure of geographical results, as the contributions of the major chemical producing regions in 2020 decrease from 14%, 29%, and 19% to 9%, 19%, and 7% for Europe, Northeast Asia, and North America, respectively. Conversely, annualised costs in the sub-Saharan Africa and SAARC regions increase substantially from 1% and 7% in 2020 to 18% and 23% in 2050, respectively. Such a rapid growth of chemical production in the regions of the Global South suggests a massive ramping of sustainable chemical production technologies as well as renewable electricity, especially solar PV, and water electrolysers, as

indicated by highly competitive solar hydrogen.¹⁰⁰ Regional annualised costs are shown in Fig. S32–S40 (ESI[†]).

LCOC at a global level, as shown in Fig. 28, shows a similar trend as the annualised costs, as the total cost per primary chemical increases globally from 900 \notin /t to 1617, 1409, and 1272 \notin /t in 2050 for the BAU, NZE 5050H, and NZE 2050L scenarios, respectively. On a regional basis, there is a noticeable deviation of LCOC in the NZE 2050 scenarios as the LCOC ranges from 1259 to 1473 \notin /t and 1157 to 1352 \notin /t in the NZE 2050H and NZE 2050L scenarios, respectively. Despite having among the lowest LCOE, the SSA and SAARC regions have among the highest levelised costs in 2050 for the NZE 2050 scenarios, compared to MENA, which has the third lowest, due to higher demands of primary chemicals. Interestingly, North America has the lowest LCOC in 2050, pointing to optimal conditions of low LCOE and high availability of plastic waste for secondary plastics.

4. Discussion

The discussion section is separated into four sub-sections. Section 4.1 will discuss the main findings of this research in **BAU Annualised Chemical Industry**

BAU Annualised Chemical Industry



Fig. 27 Annualised costs from 2020 to 2050 by cost component and by major region for the BAU (left), NZE 2050H (middle), and NZE 2050L (right) scenarios. Positive GHG emission costs in 2050 in the NZE 2050 scenarios result from the incineration of fossil plastic waste.



Fig. 28 Levelised cost of chemicals (LCOC) by cost component (top) and per major region (bottom) for the BAU (left), NZE 2050H (middle), and NZE 2050L (bottom) scenario from 2020 to 2050. Positive GHG emissions costs in 2050 in the NZE 2050 scenarios result from incineration of fossil plastic waste.

the context of other similar research. Section 4.2 provides an outlook of the green chemical industry in the context of energy system modelling. Section 4.3 considers the impacts of the defossilised chemical industry on sustainable feedstocks, and Section 4.4 discusses limitations and opportunities for further research.

4.1. Main findings

The results of this research suggest that the full defossilisation of the global chemical industry implies massive renewable electricity, green e-hydrogen, and green e-methanol demands. Across the NZE emissions scenarios studied, total electricity,

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e-hydrogen, and e-methanol demands in 2050 range from 28 909 to 40 162 TWhel, 16 069 to 23 116 TWh_{H2,LHV}, and 13 721 to 17750 TWh_{MeOH,LHV}, respectively. Such high quantities of sustainable feedstocks are confirmed by the literature, as research investigating net-zero emissions finds a total electricity demand of around 12 3074-34 70033 TWhel, the majority of which would be used for green e-hydrogen production. Although increased recycling levels can reduce the quantity of primary feedstocks, secondary plastics are not sufficient to cover global plastic demands. In 2050, secondary plastics can cover 26-31% of total plastics, and by 2100, this share increases to 51%. Saygin and Gielen⁴ found that by 2050, 42% of the plastic demand can be satisfied by increased recycling, though this also comes with demand reduction measures. Outside of chemicals-toplastics, chemicals-to-pesticides increase steadily from 24.8 TWh_{th} in 2020 to 57.3 TWh_{th} in 2050 and 82.2 TWh_{th} in 2100, which roughly correspond to feedstock demands of 55, 129, and 185 TWh_{th} in 2020, 2050, and 2100, respectively, considering a feedstock demand of 2.25 $MWh_{th}/MWh_{th,pesticide}$.¹⁰¹

Results additionally suggest that CO₂ demands by the global chemical industry will require 3.39-4.74 GtCO₂ in 2050. Research investigating a CCU-based chemical industry in 2030^{32} similarly found a global CO₂ demand of 3.72 GtCO₂, though this scenario assumes a power-to-methane-to-ammonia route rather than the power-to-hydrogen-to-ammonia route considered in this study. While this research assumes a supply of CO₂ from DAC, there are key sustainable or unavoidable point source emissions from cement production, pulp and paper mills, and waste incineration, and this potential has been estimated to be 2471 MtCO₂ in 2050.⁶² Additionally, pointsource capture of fossil plastic emissions in waste incinerators could provide CCU opportunities for the fossil carbon embedded in fossil plastics; however, this potential is limited to only several hundred MtCO₂. For the global CO₂ demands of this research, the total point source emission potential could satisfy 52-74% of total CO2 demands by 2050, which could increase the economic viability of power-to-chemicals. This may also lead to co-location of e-chemical production with low-cost point source CO₂ emissions.¹⁰²

The geographical results suggest that the most rapid growth of green feedstocks for most regions will occur from 2030 to 2050, with regions that do not reach their peaks until the end of the century only having moderate growth in feedstock demands. The most notable exception to this trend, though, can be seen in the results for sub-Saharan Africa, as the largest growth in demand occurs in the latter half of the century. While today's chemical industry is largely dominated by countries of the Global North, this structure will likely shift to have higher shares of chemical production in the countries of the Global South, which has already been observed in net-zero emissions scenarios for the steel industry.¹⁰³ Research for the chemical industry in Europe finds that in a power-to-X dominated structure, the total indirect electricity demand for chemicals reaches 2634 TWh_{el}⁴³ which confirms the indirect electricity demand for Europe in this study of 2609 TWhel in the NZE 2050L scenario. Additionally, e-plastics production alone in

Europe could result in a total electricity demand of 1615 $TWh_{\rm el}.^{104}$

Analysis of annualised and levelised cost structure finds that the NZE 2050L scenario, with the highest shares of e-chemicals, has the lowest annualised costs by 2050, pointing to the viability power-to-X economy¹⁰⁵ basis for the global chemical industry. Such levels of renewable electricity supply are feasible, as research in the 100% RE system analyses has found a cost-optimal renewable electricity generation of over 100 000 TWhel from solar PV and wind power alone.¹⁰⁶ While this research finds that a high electrification of the global chemical industry leads to the lowest annualised costs, Zibunas et al.³³ found that higher biomass shares, in conjunction with increased recycling and some electrification, lead to lowest annualised costs among net-zero emissions scenarios compared to the high electrification scenario. This contrasting finding is heavily influenced by the electricity costs used, especially for water electrolysis, which ranges from 10.0 to 18.8 €/MWhel in this research, strongly driven by the low-cost solar PV. Additionally, relevant to the economic viability of the green chemical transition is the use of GHG emissions pricing mechanisms, as the BAU scenario without GHG emissions costs would be the least cost scenario at 1468 b€ in 2050 with a LCOC of 1044 €/t. Furthermore, sustainable chemical production during the transition would consistently be more expensive than fossil routes, especially for HVCs that have low feedstock emissions. Therefore, some additional economic incentives may be required for green chemical production, such as a carbon credit for the CO₂ used in e-chemicals.¹⁰⁷

4.2. Green chemical industry in energy system modelling

While the energy transition of energy systems to high shares of renewables is well understood,^{106,108} the global transition of non-energy feedstocks to 100% renewable feedstock has not yet been investigated in an energy system model. However, it has been conceptionally added to the first energy system model,¹⁰⁹ and it was applied for Europe,¹¹⁰ which is the only known study applying all required e-fuels and e-chemical buildings blocks for the sustainable energy-industry system transition.⁶² The chemical sector alone is responsible for over 11000 TWh_{th} of fossil fuel consumption, and the share of chemical production in total fossil fuel demands is expected to increase as other sectors defossilise.¹ Furthermore, similar to other industry sectors, high electrification of production processes is not sufficient to eliminate CO2 emissions. The chemical industry is among the most difficult to abate sectors because of the requirement of hydrocarbon feedstocks and the high costs of e-chemicals. The most feedstock emission intensive processes in the chemical industry are ammonia and methanol production; thus research has largely focused on the defossilisation of these processes through power-to-X^{13,19,63,64,70,111-116} and biomass^{85,117-121} routes. While methanol-to-chemicals routes have gained increased attention in the literature for methanolto-olefins^{24,72,74,75,122} and methanol-to-aromatics,^{76,77,123} much of this research has focused on coal-based methanol rather than power-to-methanol.

Although feedstock switching is the key to defossilisation of the chemical industry, it is also crucial to understand how feedstock quantities can change because of process switching. Taking ethylene, this research shows a global transition from steam crackers based on oil products to MTO. Outside of the main products, the by-product structure of these routes is noticeably different, and research suggests that some steam crackers may be necessary to satisfy global butadiene demands. which today is covered completely by steam cracker byproducts.⁴ Furthermore, the feedstock requirements per ton of chemicals can vary significantly by the process and feedstock, which is especially true for the MTA process, which has a BTX yield of only 16%.26 Therefore, a one-to-one feedstock switching for modelling of global chemical demands may not be appropriate, and more detailed examination of feedstock requirements for the primary platform chemicals is necessary, as performed in this research. The approach presented in this research can thus provide scalable sustainable pathways for the global chemical industry that fully phases out fossil fuels that can be applied in energy system models.

Understanding the transition of the global chemical industry towards sustainable feedstocks in conjunction with the larger global energy-industry system transition is still lacking.¹⁰⁶ The 2022 IEA World Energy Outlook's NZE 2050 scenario⁹⁹ still requires over 20 EJ of fossil feedstocks for chemical production, with electrification routes only being utilised for e-ammonia and e-methanol production. However, when considering the quantities of feedstocks that would need to be defossilised, a full consideration of chemical feedstocks is extremely relevant to the complete energy-industry transition perspective. When examining the NZE2050 scenarios, the total feedstocks required increase from 11 029 TWhth of fossil feedstocks in 2020 to 20628 TWhth of e-chemical feedstocks in 2050, and 25515 TWh_{th} in 2100. Fully defossilised feedstock demands would then require primary electricity and bioenergy demands of 27759-33 082 TWhel and 2548-9848 TWhth in 2050, and 31 026-42 108 TWhel and 2948-16846 TWhth in 2100, respectively. Therefore, detailed estimates of the primary energy and final energy demands of the chemical industry, as performed in Keiner *et al.*,³⁸ are essential for developing comprehensive perspectives on the energy-industry transition, as it may be the most relevant in terms of primary and final energy demands for all industrial sectors.

Insights into the European energy-industry transition¹¹⁰ are among the first to provide defossilisation pathways for chemical feedstocks, and point to an interesting dynamic where e-naphtha, a by-product from the Fischer-Tropsch process for e-fuels production,¹²⁴ can be used as a feedstock in the chemical industry. Given the low yield of BTX aromatics from the MTA process, e-naphtha by-products may be best utilised in the catalytic reforming of naphtha, which has a much higher aromatic yield. Ikähemo *et al.*¹⁷ investigated the interactions of e-ammonia production in the Northern European energy system and found that e-ammonia could act as a form of long-term energy storage and that waste heat from the power-to-ammonia processes. Given that both the MTO and MTA processes are

exothermic,⁷⁴ with excess heat from these processes alone reaching over 861–1134 TWh_{th} by 2050, there may be opportunities for analogous usage of waste heat both in other industrial processes of district heating systems, and as valuable heat for reducing CO₂ sourcing cost in DAC units.¹²⁵ Additionally, the high wastewater by-product from MTO and MTA could supply the high water demands required for electrolysis, which may be extremely relevant if there is a gradual shift of the global chemical industry towards production in regions with low-cost electricity and large amounts of the available land. CCU for the chemical industry could increase the circularity of the global chemical industry in CO₂, energy integration, and water.⁴⁵ However, the power-to-methanol and methanol-to-chemical production sites may be part of split value chains, which would reduce the potential to utilise this wastewater by-product.

4.3. Impact of the global chemical industry on sustainable feedstock demands

Since the NZE 2050L scenario emerges as the least cost scenario studied, many regions of the world may opt for an e-chemical production landscape, especially those with limited sustainable biomass resources. Of the industrial sectors, the chemical industry stands to become the industry with the highest demand for e-hydrogen, as this research finds e-hydrogen demands in the range of 16069-23116 TWh_{H2,LHV}. In a chemical sector completely supplied by e-ammonia and e-methanol, the total e-hydrogen demand could similarly be within the range of 19000 TWh_{H2,LHV}.¹² Furthermore, if all chemical demands in 2030 were to be supplied by e-methanol and e-methane for ammonia, total e-hydrogen demands would total 19831 TWh_{H2,LHV}.³² This e-hydrogen demand may be larger than the next two largest industrial consumers, the cement and steel industry, whose e-hydrogen demands may range between 4900 and 10400 TWh_{H2,LHV}¹²⁶ and 2800 and 4400 TWh_{H2,LHV},¹⁰³ respectively. In the NZE 2050 scenario studied, the total installed electrolyser capacity would reach 5.2-6.2 TW_{H2,LHV} by 2050, assuming flexible operation according to renewable electricity generation profiles,³⁶ which represents about 31–36% of the total estimated about 17 TW_{H2,LHV} of the electrolyser demand by 2050.100 The steel and cement industries then may require additional terawatt-scale capacities of both electrolysers and solar PV and wind power to ensure low e-hydrogen production costs.

The solar PV capacity in 2021 represented 50% of the global added power generation capacity.¹²⁷ It is ramped at a high CAGR of 20–30%/a in the long term and a silicon production capacity of more than 900 GW/a is forecasted by 2050,¹²⁸ while the growth is projected to continue.^{36,129} Studies have found that 70–80% of the total global electricity generation could be contributed by solar PV by 2050,³⁶ which is a consequence of low-costs, the historic fastest growth and phase-in of any energy source,¹³⁰ and the PV industry that is capable to continuously grow faster in manufacturing capacities than the demand.^{129,131} It is estimated that the total installed PV capacity by 2050 could be up to 70 TW,¹²⁹ of which 18.6 TW (27%) would be required for the chemical industry, assuming 76% of the

40 382 TWh of the electricity demand for the NZE 2050L supplied by the solar PV for a global average yield of 1650 kWh/kWp.³⁶ Furthermore, CO_2 capture facilities, either a point source or DAC, will need to be in the order of GtCO₂/a, as this research finds a total CO_2 demand of 3.34–4.74 GtCO₂/a.

During the transitionary period, biomass could be used to reduce chemical industry emissions without such a large electrolyser and CO_2 capture plants, though there would still be the need for megaton-scale bio-ammonia and bio-methanol plants. In this research, biomass resources were prioritised for bioplastic chemicals, which, with limited shares of bio-ammonia and direct bio-methanol, require 16 799 TWh_{th} of biomass by 2100, corresponding to 60% of the global sustainable biomass resource of 100 EJ⁹⁴ (27 800 TWh). While the share of biochemicals could be higher, sustainable biomass may also be in competition with use in other energy sectors, such as biofuels for transport, bioenergy for heat supply, and compete with food production.⁴² Without increases in plastic recycling, total biomass consumption for chemicals could reach up to 101 EJ,³³ just about exceeding sustainable limits.

Bio-based plastics are among the chemical products that have received the most attention in recent years;¹ however, a challenge remains with the biodegradability of bio-plastics, as conventional bio-based plastics, e.g., bio-based polyethylene and polyethylene terephthalate, are not biodegradable nor compostable,¹ though there is research to increase the biodegradability of conventional plastics.¹³² Furthermore, life-cycle analysis results suggest that a power-to-polyethylene route may be more beneficial in terms of climate and biodiversity impacts compared to the bio-polyethylene route.133 The use of limited biomass resources may be best prioritised in the production of biodegradable bioplastics such as polybutylene adipate terephthalate and polylactic acid, though responsible end-of-life practices are still necessary for biodegradable plastics.¹³⁴ The co-evolution and integration of e-chemicals and bio-chemicals may be a viable pathway forward, as low-cost electricity could be coupled with biogenic carbon inputs,104 along with the parallel development of biodegradable plastics.

4.4. Limitations and future work recommendations

Due to the complexity of the global chemical industry, not all data regarding chemical flows were readily available, and a lack of open access to production data in the chemical industry has been highlighted by Levi and Cullen.8 Therefore, the model developed here is not able to cover every relevant chemical production process, but major routes, especially those for ammonia, methanol, and the HVCs, are fully covered. Additionally, data for the chemical industry is typically provided at a global level; thus regional distribution is based on feedstock usage, which may not perfectly capture where chemical products are finally used, as chemicals imports and exports are not transparent. The regional production and use landscape should be placed under further scrutiny in a transition to a green chemicals landscape, especially one dominated by e-chemicals. Previous research for e-steel has suggested that imports of intermediate iron and final steel from regions with abundant solar resources, many of which are in the Global South, may be

lower in cost than domestic e-steel.^{135–137} Chemical production may follow a similar trend where chemical plants are sited in regions with the best RE resource availability. For China, the United States, and the Middle East, there may be sufficiently available RE resources to continue high levels of domestic production; however, Europe may have challenges with producing economically competitive e-chemicals compared to the best sites in the world. The solution for Europe may be to largely import e-methanol as a new bulk chemical so that HVC can be produced, similarly to the use of imported fossil feedstocks as of today. Therefore, additional research should investigate new supply chain configurations that can best capitalise on low-cost solar electricity to increase the economic viability of e-chemicals, especially for e-olefins and e-aromatics.

When considering the economic viability of e-chemicals compared to fossil-based and biomass-based chemicals, electricity costs are the most relevant factor as primary electricity is the key input for e-chemicals. While the indirect LCOE applied to electrolysers, which do not need to match specific load profiles, has low uncertainty when supplied by hybrid solar PV-wind power plants, the direct LCOE applied to the chemical production plants, which are assumed to operate under near baseload conditions, has higher uncertainty. The direct LCOE applied in this study, based on the transition of the entire energy system,³⁶ may better match the legacy cost of existing systems, and thus lead to higher LCOE. Comparatively, the approach taken by Fasihi and Breyer⁹⁷ would be better adapted to near baseload supply of the chemical industry and new greenfield investments based on RE supply. In terms of annualised costs, applying the latter LCOE methodology would lead to increased annualised costs of 6.7% in 2020, but a decrease of annualised costs of 66 b€, or 3.0%, in 2050 in the NZE 2050L scenario. Additionally, the emission reductions from applying direct and indirect LCOEs based on greenfield hybrid solar PV-wind electricity supplies are 236, 110, and 56 MtCO_{2eq} in 2020, 2030, and 2040, respectively, in the NZE 2050L scenario.

There is also high uncertainty with maximum potential recycling rates for plastics, as recycling rates in 2050 can range in the literature from 54¹ to 66⁴%, and a recycling rate of 60% was established for the NZE year in this study. Potentially higher plastic recycling rates may be possible, which would reduce the primary chemical demand and thus the sustainable feedstock demands. Nevertheless, plastic chemicals comprise around 61% of the total chemical demand modelled in this research in 2050; therefore, significant quantities of sustainable feedstocks will still be required for non-plastic chemicals. Furthermore, as mentioned in Section 4.2, e-naphtha by-products from Fischer-Tropsch plants may be utilised to substitute green methanol demands,¹²⁴ though quantities of e-naphtha available require comprehensive energy-industry system analysis and was thus not considered in this research.

5. Conclusions

The defossilisation of the global chemical industry implies significant e-hydrogen and renewable electricity demands as

green ammonia and green methanol compose the background of a sustainable chemical industry. This research provides the first pathways to a sustainable chemical industry to the end of the century with a high global-local resolution of 145 LUT regions. While annualised costs for a green chemical industry can be lower than business-as-usual conditions, certain requirements must be met, such as

• Gigaton-scale CO_2 as the raw material is required either from point sources, *e.g.*, cement, waste incinerators, or pulp and paper industries, or from direct air capture. Furthermore, terawatt-scale solar PV and wind power as well as electrolysers will be required for the chemical industry alone. The wastewater by-product of the methanol-to-olefins and methanol-toaromatics processes may be usable by water electrolysers to reduce the freshwater demand of the chemical industry if electrolysers are co-located with chemical production facilities, which would be of high relevance in dry regions with excellent solar resources.

• High shares of plastic recycling are required to reduce the total feedstock requirement for primary chemicals, but the available secondary plastics are not sufficient to completely substitute primary plastic production.

• High biomass routes lead to higher annualised costs compared to high shares of e-chemicals, suggesting that limited sustainable biomass resources may be better utilised in emerging biodegradable bioplastic markets, or as a source of biogenic CO_2 coupled with e-chemical production. Furthermore, use of first-generation biomass sources should be avoided due to land-use constraints and competition with food production.

• A full transition of the global energy system towards absolute zero CO_2 emissions is required to completely decouple fossil fuels from the chemical industry. Without a full defossilisation of the energy system, increased shares of e-chemical production could lead to an increase in CO_2 emissions from the chemical industry due to the high quantities of electricity required. Furthermore, this research finds that all process demands, both direct electric and heating, can be met by direct electricity, electric heating, and heat pumps.

Global feedstock demands for the chemical industry are expected to increase to the end of the century, with the highest growth being experienced in the emerging economies of the Global South, with global green ammonia and methanol demands reaching 2365 TWh_{NH3,LHV} and 23 343 TWh_{MeOH,LHV} in 2100, respectively. This global growth in feedstock demands is driven primarily by sub-Saharan Africa and South Asia. Regional development of the chemical industry transition may affect feedstock choices for green chemicals, and new chemical production hubs may emerge in regions with high availability of land and low-cost renewable electricity, especially from solar photovoltaics. Access to sustainable feedstocks may be the largest bottleneck hindering green chemical production, thus improved waste management and higher recycling rates can serve to reduce primary feedstock demands. Therefore, rapid defossilisation of global energy systems through low-cost solar electricity, along with improved plastic recycling, can catalyse an economically viable defossilisation of the global chemical industry.

ASU	Air separation unit
BECCS	Bioenergy carbon capture and storage
bio-Ammonia	Biomass-based ammonia
bio-Methanol	Biomass-based methanol
BNL	Belgium and the Netherlands
BTX	Benzene, toluene, and xylenes
CAGR	Compound annual growth rate
CCS	Carbon capture and storage
CCU	Carbon capture and utilisation
CR	Catalytic reforming of naphtha
crf	Capital recovery factor
DAC	Direct air capture
DCC	Deep catalytic cracking
e-Ammonia	Electricity-based ammonia
e-Hydrogen	Electricity-based hydrogen
e-Methanol	Electricity-based methanol
GHG	Greenhouse gas
HVC	High value chemical
IAM	Integrated assessment model
IEA	International energy agency
IPCC	Intergovernmental panel on climate change
LCOC	Levelised cost of chemicals
LCOE	Levelised cost of electricity
LPG	Liquefied petroleum gas
MENA	Middle east and north Africa
MeOH	Methanol
МТО	Methanol-to-olefins
MTA	Methanol-to-aromatics
NZE	Net-zero emissions
OECD	Organisation for economic co-operation and
	development
OPP	On-purpose propylene
RE	Renewable energy
TDP	Toluene disproportionation
THD	Toluene hydrodealkylation
TRL	Technology readiness level
SSA	Sub-Saharan Africa
SAARC	Southeast Asia association for regional
	cooperation
WACC	Weighted average cost of capital

Conflicts of interest

There are no conflicts to declare.

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