Green Chemistry



PAPER

View Article Online



Cite this: *Green Chem.*, 2022, **24**, 6671

Making more from bio-based platforms: life cycle assessment and techno-economic analysis of *N*-vinyl-2-pyrrolidone from succinic acid†

Moritz O. Haus, (1) ‡ Benedikt Winter, (1) ‡ b,c Lorenz Fleitmann, (1) b,c Regina Palkovits (1) ** and André Bardow (1) ** b,c,d

The prospective assessment of the environmental and economic costs of emergent technologies based on biomass feedstocks has a vital role in guiding academic research and corporate investments. This study evaluates a new pathway for the synthesis of the established monomer N-vinyl-2-pyrrolidone (NVP) from succinic acid, a suggested biorefinery platform chemical. For this purpose, conceptual process design is combined with recent progress in catalysis research to deliver the life cycle inventories of the novel process route. LCA projects that bio-based NVP production reduces global warming impacts by 25-53% compared to the fossil alternative practiced today. Within this range, the magnitude of reductions depends on the biomass feedstock, the chosen hydrogenation catalyst, and separation technology. However, these global warming impact reductions have to be weighed against increasing other environmental impacts, mainly eutrophication and acidification, which are associated with farming and the expenditure of noble metal catalysts. The operational cost analysis suggests that succinic acid-based NVP production is competitive if the substrate cost ratio (succinic acid/ γ -butyrolactone) is lower than 0.7. The insights gained in this study provide goals for catalyst research and process development.

Received 30th March 2022, Accepted 8th August 2022 DOI: 10.1039/d2gc01219g

rsc.li/greenchem

Introduction

The integration of biomass feedstocks into traditionally fossil-based production sectors may be key to enabling human prosperity within the resource and climate constraints of planet earth. The biomass-based (short: bio-based) synthesis of chemicals and fuels has therefore been an important subject of political efforts, 5-8 societal discussions, and academic

^aChair of Heterogeneous Catalysis and Technical Chemistry, RWTH Aachen University, Worringerweg 2, DE-52074 Aachen, Germany.

E-mail: palkovits@itmc.rwth-aachen.de

‡These authors contributed equally to this work.

research. ⁹⁻¹⁴ In this context, polylactic acid, ^{15,16} bio-ethanol ¹⁷ and bio-diesel ¹⁸ have emerged as encouraging examples, showing that bio-based production can be brought to the commercial scale.

Yet, a broader transition to biomass feedstocks requires biorefineries that integrate a large network of processes converting the highly functionalized substrates into energy carriers and chemical building blocks. Pegarding chemical building blocks, several studies have evaluated potential bio-economy platform chemicals based on criteria such as application versatility, market potential, and technology readiness. Often-reported key candidates include chemicals of carbohydrate origin, such as hydroxymethylfurfural (or derived furandicarboxylic acid), sorbitol (or derived glycols), and succinic acid. Consequently, value chains starting from these compounds should be of special interest to the chemical research community.

In the specific case of succinic acid, a major research focus has been on liquid-phase hydrogenations yielding γ -butyrolactone (GBL), 1,4-butanediol, and tetrahydrofuran (THF). While each of these products can be preferentially formed with the correct choice of catalyst system and reaction conditions, some selectivity issues remain due to the sequential nature of the ongoing hydrogenation steps. More importantly, however, current succinic acid price estimates (\sim 2.5 \$

^bEnergy and Process Systems Engineering, ETH Zurich, Tannenstraße 3CH-8092 Zurich, Switzerland. E-mail: abardow@ethz.ch

^cInstitute of Technical Thermodynamics, RWTH Aachen University, Schinkelstraße 8, DE-52062 Aachen, Germany

^dInstitute of Energy and Climate Research (IEK 10), Research Center Jülich GmbH, Germany

[†] Electronic supplementary information (ESI) available: Contents: 1 Experimental laboratory procedures; 2. Experimental results on amidation-hydrogenation; 3. Life cycle inventories; 4. Life cycle impact assessment; 5. Process descriptions; 6. Estimation of thermodynamic properties; 7. Operational cost analysis; 8. Exemplary comparison of unit operations; 9. Analysis of alternative scenarios for fossil-based NVP production; 10. Detailed waste stream compositions and LCIs; 11. Tabulated LCA results. See DOI: https://doi.org/10.1039/d2gc01219g

per kg)³¹ will challenge the prospects for economical production of comparably inexpensive lactones, alcohols and ethers (*e.g.* 1,4-butanediol price of 2.3 \$ per kg).³²

In contrast, the synthesis of heteroatom-containing chemicals from biomass may allow for easier market access due to higher prices and long synthesis pathways starting from fossil resources. 1,33 Concerning the conversion of succinic acid into such chemicals, pyrrolidones could offer an attractive option with applications e.g. in pharmaceuticals and specialty polymers. As a potential pathway, White et al. discussed producing N-methyl-2-pyrrolidone (NMP) and 2-pyrrolidone (2PYD) from succinate-containing fermentation broth.³⁴ The respective process comprises thermal imide formation, distillation, and reduction over a rhodium catalyst (220 °C, 100 bar H2). While the catalytic conversion of succinic anhydride into N-phenyl-2pyrrolidone was also reported,35 the total number of studies on value chains from succinic acid to pyrrolidones has remained comparably limited, with patents comprising the bulk of available literature. 36,37

In this context, we recently proposed a sequence of liquid phase amidation–hydrogenation and gas-phase dehydration processes to synthesize *N*-vinyl-2-pyrrolidone (NVP) from aqueous succinic acid solutions (Fig. 1).^{39,40} Yields of 70 mol% (succininc acid to NVP) were achieved with commercial hydrogenation catalysts and improved by further material development.⁴¹ Consequently, this reaction sequence seems to be a promising way of obtaining a high-value monomer from a potential biomass platform using two conversion steps. While this value proposition appears intriguing, a detailed evaluation of process inputs and outputs is required to determine the expected merit of the proposed transformation.

Here, life cycle assessment (LCA)⁴² offers a tool for the holistic analysis of value chains, as already demonstrated for biomass utilization scenarios. ^{11,43} Previous early-stage LCA studies have resolved the influence of design choices, including catalyst materials and biomass feedstocks, thus helping to guide future research priorities. ^{44,45} Consequently, to quantify the environmental and economic impact of bio-based NVP production, an LCA study on the current state of the succinic acid-to-NVP conversion technology is presented herein.

The study starts with defining the underlying LCA methodology (section 2), including the goal and scope definition (section 2.1), the system boundaries for bio- and fossil-based NVP production (section 2.2), the methodology for compilation

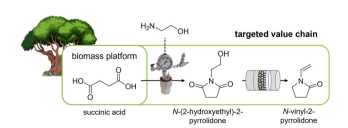


Fig. 1 Targeted value chain from succinic acid, as a bio-based platform chemical, to N-vinyl-2-pyrrolidone.³⁸

and evaluation of process inventories (section 2.3), process simulations for bio-based NVP production (section 2.4) and economic evaluation (section 2.5). Life-cycle inventories of different process options are presented and discussed next (section 3.1). The following life cycle impact assessment (section 3.2) focuses first on global warming impact and is extended to other non-tox impacts of the CML2001/ 2016 method. 46 Toxicology related impacts are available in the ESI.† Next to these environmental metrics, operational cost estimates are presented as an economic indicator (section 3.3). The comparative analysis highlights the environmental and economic potential of replacing fossil NVP production, considering different biomass feedstocks, separation technologies, and succinic acid market scenarios. After discussing uncertainties in the early stage evaluation (section 3.4), directions for future catalyst and process development are formulated (section 3.5). To the best of our knowledge, we present the first LCA study of bio-based pyrrolidone production.

2. Methodology

2.1. Goal and scope definition

The presented LCA aims to compare the environmental impacts of 1 kg of NVP produced through conventional and succinic acid-based processes. Thus, the functional unit of the evaluation is 1 kg_{NVP} (molar purity >99%). Moreover, cradle-togate system boundaries are applied – including all processes from feedstock extraction to the factory gate. Since conventional and bio-based NVP are chemically identical at the factory gate, the use-phase of NVP can be assumed to be independent of the chemical's origin. Consequently, gate-to-grave impacts on LCA results cancel in the comparative approach and are therefore not considered. 47,48

Moreover, the evaluation is prospective and relates to a potential implementation of NVP production from succinic acid, obtained by sugar fermentation. To assess the impact of technological choices and developments that cannot be fully foreseen at this stage, several catalyst and separation scenarios are included. Furthermore, the source and type of biomass feedstock has been shown to have a vast effect on the environmental impact of bio-based chemicals. 45,49 Thus, we include a set of typical first and second-generation feedstocks for sugar production in this study: sugar beet (base case), sugar cane, corn, and corn stover. While these feedstocks are often associated with distinct production regions,50 other regional differences are not considered for the early-stage assessment, and production of succinic acid and pyrrolidone is assumed to occur in Europe. Finally, the purified C5/C6-sugars derived from the named biomass feedstocks are assumed to yield identical results in succinic-acid production, though further research on feedstock variability may be required. 51-53

2.2. Process description and system boundaries

2.2.1. Fossil-based production. Within this analysis, conventional (*i.e.* fossil) NVP production is based on the maleic

anhydride platform. This route is assumed to be the most economical way of accessing the required GBL intermediate, though older alternatives based on acetylene and formal-dehyde (so-called Reppe chemistry) remain in operation.⁵⁴ First, maleic anhydride is converted to GBL by selective gasphase hydrogenation over base metal catalysts.^{34,54} Subsequently, the product is aminated to 2-pyrrolidone, which in turn can be reacted with acetylene to yield NVP (Fig. 2).⁵⁵

Several process options are disclosed in the patent literature, especially for realizing the last two stages (from GBL to NVP). ⁵⁶⁻⁶⁰ A comparison of these options suggests that high-pressure, liquid-phase amination in a concentrated, aqueous solution without catalyst is state-of-the-art in 2-pyrrolidone synthesis. ⁵⁶ Subsequently, vinylation is performed with pressurized acetylene using potassium pyrrolidate, which forms from 2-pyrrolidone and KOH in water-free conditions, as a homogeneous catalyst. ^{57,58}

While this option is not implemented herein, an extensive array of patents has targeted the avoidance of acetylene in generating the vinyl-group of NVP. 61,62 Most notably, the condensation of GBL and monoethanolamine yields *N*-(2-hydroxyethyl)-2-pyrrolidone (HEP), which can be dehydrated to NVP. 63 Despite its dependence on fossil carbon due to the commercial sources of GBL, this alternative has inspired part of the succinic acid-based NVP production chain. This option also opens a path to bio-based NVP from bio-based GBL, which is not within the scope of the current article. 64

2.2.2. Bio-based production. In the bio-based value chain, low-pH yeast fermentation of sugars yields succinic acid, which is purified by direct crystallization. The acid is subsequently mixed with monoethanolamine in a concentrated aqueous solution. High-pressure, liquid-phase hydrogenation of the mixture yields HEP as main product, which is then dehydrated in a gas-phase reaction, using nitrogen as an inert

diluent (Fig. 3).³⁹ Additional details on process layouts and conditions are given in the process simulation section 2.4.

2.3. LCI compilation and LCIA

2.3.1. Life cycle inventories (LCI). The life cycle inventory of this study is based on various data sources to best reflect the current state of knowledge. Where possible, data from the GaBi LCA database (V8.7 SP 40)⁶⁶ and from published literature was preferred. Secondly, inventory data was obtained from the IHS Markit chemical economics handbook.⁶⁷ If no data was available in either of these sources, process simulations based on available patents or novel process ideas were used to generate the lacking inventories.

Since carbohydrates are frequently used as starting point of biomass-based chemical production, inventory data for the required agricultural and processing steps is present in scientific literature. The evaluation at hand specifically uses data compiled by Winter *et al.* for bio-based aniline production (ESI, section 3†).⁵⁰ Moreover, information on fermentation to yield succinic acid was taken from Cok *et al.*⁶⁵ Lastly, commercial inventory data on ethanolamine produced from ethylene oxide was obtained from IHS Markit.

In the case of fossil production, maleic anhydride reduction to GBL is the first process stage absent from the LCA database. Thus, commercial inventory data was obtained from IHS Markit. Subsequent steps in both value chains – starting from GBL and succinic acid, respectively – had to be analyzed by process flowsheet simulations due to a lack of published inventory data. These flowsheet simulations yielded a variety of output streams, some of which require further treatment. The output streams were categorized as (i) product, (ii) inert gas purge, (iii) wastewater, (iv) organic residue, and (v) combustible gas purge. While inert gases (ii) are released to the environment without further treatment, wastewater (iii) is dis-

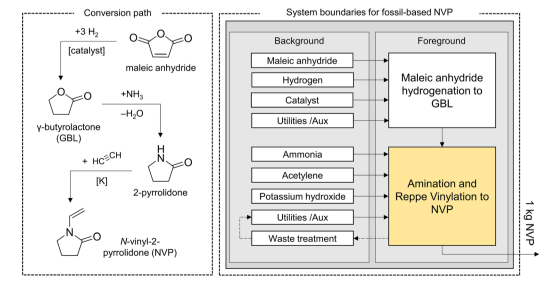


Fig. 2 Conversion path and system boundaries for the evaluation of NVP synthesis from the maleic anhydride platform. Yellow color highlights process simulations conducted in this work.

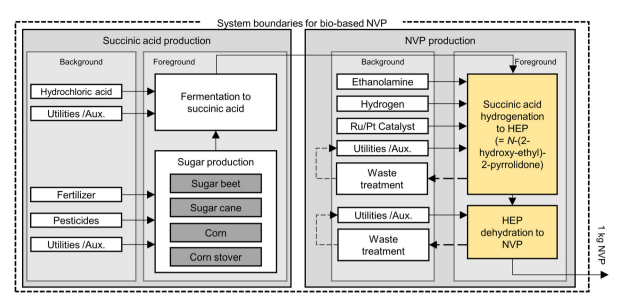


Fig. 3 System boundaries for the evaluation of NVP synthesis from biomass *via* succinic acid. Yellow color highlights process simulations conducted in this work. Refer to Fig. 1 for a chemical representation of conversion steps.

carded through a sequence of wet air oxidation and mechanical-biological treatment, and organic residue and combustible gases (iv-v) are jointly incinerated. The required inventory data for the treatment of (iii-v) is based on the literature analysis of waste treatment facilities in commercial plants. ^{68,69}

2.3.2. Life cycle impact assessment (LCIA). The compiled inventories of NVP production scenarios were evaluated using the CML2001/2016 method due to its wide acceptance. Calculations were run in Gabi V8.7 SP40 using the list of background processes provided in the ESI (section 4a†).

For the treatment of biogenic carbon, we followed Pawelzik et al. in accounting for all its uptake from and release into the atmosphere.⁷⁰ Consequently, the GWI of bio-based NVP explicitly accounts for the amount of carbon uptake in plant cultivation but also the direct release during processing (e.g. fermentation) and NVP production. This treatment reflects the carbon flows from cradle-to-gate in line with the goal and scope of this study. Assuming combustion as an exemplary end-of-life scenario, the expansion towards cradle-to-grave boundaries closes the circle of carbon capture and release in the case of bio-based NVP, whereas conventional NVP would be burdened with additional carbon emissions, ultimately stemming from a fossil reservoir. However, our approach of explicitly accounting for biogenic carbon leads to very small and sometimes negative - GWI attributed to succinic acid from cradle-to-gate. This negative carbon footprint does not imply that succinic acid production is a carbon-negative technology since the comparative assessment does not include end-of-life emissions.71

Moreover, the impacts of indirect land-use change (iLUC) have been discussed as a major factor in assessing chemicals and fuels from biomass.⁷² To incorporate this aspect despite the remaining large uncertainty in this field,⁷³ iLUC effects on global warming impacts are estimated explicitly by exploiting

knowledge on bio-ethanol which shares the sugar basis with bio-based NVP. The variability of results related to iLUC is shown by adapting values from several studies for each of the discussed biomass feedstocks. Lastly, it is noted that the models applied in the original iLUC research papers often do not exclude effects from direct land-use change. Keeping with the authors of those studies, the adaptation of their results to the bio-based NVP scenario is still referred to as iLUC. The reader is referred to the supplementary (section 4) for additional details on iLUC and other LCA aspects.

2.4. Process simulations

As discussed in section 2.3, part of the LCI data is obtained by rigorous process simulations using the flowsheeting software Aspen Plus® V11. Required process information is derived from scientific and patent literature as well as additional experiments (ESI, section 1 and 2†). In particular, amidationhydrogenation of succinic acid with ethanolamine (Fig. 4, $n_{\text{amine}}/n_{\text{acid}} = 1$, $m_{\text{water}}/m_{\text{acid}} = 1$) is implemented as a batch operation at 200 °C and 200 bar H2, with yield and by-product formation depending on catalyst choice. Three alternative simulations are implemented: (1) a commercial benchmark catalyst (Ru/C, 74% yield), 39 (2) a recently described optimized system (Pt-Re/TiO₂, 83% yield)⁴¹ and (3) a hypothetical future case with much reduced by-product formation (98% yield). To account for uncertainty in the handling of oligomeric hydrogenation by-products in the bio-based process, an alternative HEP separation is also considered supported by experiments (ESI, section $2b + c\dagger$): a sequence of (vacuum) distillations is implemented with or without prior removal of oligomers by extraction, resulting in conservative and optimistic process scenarios, respectively. Temperatures above the product's normal boiling point (>280 °C) are assumed to cause thermal stability issues. This limit imposes the need for vacuum separGreen Chemistry Paper

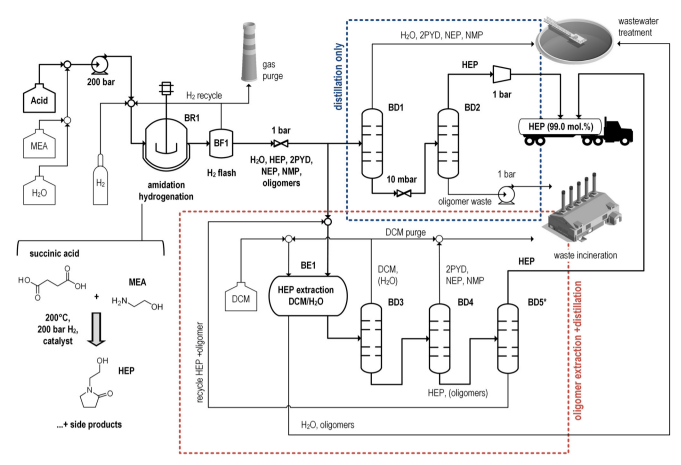


Fig. 4 Flowsheet of HEP production from succinic acid and monoethanolamine as a first stage of the *Bio*-process simulation. Downstream purification alternatives (*Bio-x-D* and *Bio-x-Ex*) are shown in the blue and red boxes. "x" serves as indicator of the three hydrogenation catalyst options discussed in the text. *separation of oligomer traces leftover from extraction; BF labels reactors, BF flashes and BD distillation columns.

ation, especially when high-boiling oligomers are removed by distillation.

Subsequent dehydration of purified HEP (Fig. 5) is based on results from continuous experimentation with a cost-

effective Na_2O/SiO_2 catalyst. ^{39,40} Conversion of 10 vol% HEP in nitrogen carrier gas at 385 °C yields 95% NVP and 3% of 2-pyrrolidone by-product. Pressure levels (199–66 mbar) in the ensuing distillation train are chosen in accordance with

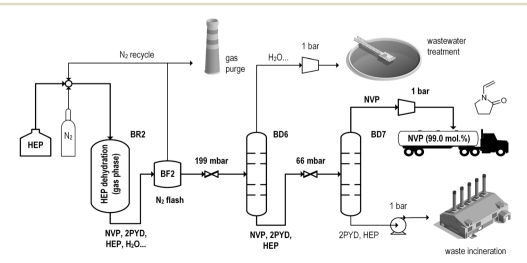


Fig. 5 Flowsheet of NVP production from HEP as a second stage of the *Bio*-process simulations.⁸²

Table 1 Overview of process scenarios analyzed in this study

Process description	Acronym
Fossil NVP production from GBL (reference) Succinic acid-based NVP production using Ru/C catalyst, distillation only Succinic acid-based NVP production using Ru/C catalyst, extraction–distillation Succinic acid-based NVP production using Pt–Re/TiO ₂ catalyst, extraction–distillation Succinic acid-based NVP production using ideally optimized catalyst, extraction–distillation	Fos Bio-Ru-D Bio-Ru-Ex Bio-Pt-Ex Bio-Id-Ex

patent-reported product stability.⁸³ The distillation train also removes co-formed water and acetaldehyde. Due to spatial limitations, flowsheets and detailed descriptions of the fossil-based process simulation are presented in the ESI (section 5†). All compared process scenarios are summarized in Table 1.

Notably, the simulated processes contain a mixture of batch and continuous operations, with flowsheets omitting intermediate storage containers and heat exchangers for the sake of clarity. Reactors, distillation columns, and continuous extractions are implemented as RStoic, RadFrac, and Extract Aspen® models. All simulations are based on Henry (permanent gases) and NRTL (other components) thermodynamic models. Most single component parameters, especially those related to thermal separation (c_p^{ig} , p_v , ΔH_v), are sourced from databases available in Aspen Plus®. For a few cases, including some properties of HEP, NVP, and oligomers, property data is estimated by COSMO-RS and the Benson group contribution method. ^{84,85} Details are given in the ESI (section 6†). NRTL parameters for key component binary interactions unavailable in Aspen® databases are also estimated by COSMO-RS.

2.5. Economic evaluation

The life-cycle inventories for the process scenarios from the environmental impact assessment allow estimating operational cost estimates as an early-stage economic assessment. Prices for chemicals and utilities are given in the ESI (section 7†). All prices are given in US dollars in the year 2020. The cost of catalysts is estimated through heuristics given in the CatCostTM calculation tool (V1.0.4). 86

Given the scope of the evaluation and the low technology readiness level of bio-based NVP, capital investment is not estimated. However, the ESI (section 8†) also contains comparative lists of the main process equipment needed for conventional and bio-based NVP production. The units are characterized by their operating conditions and product-normalized flowrates from the equilibrium-based simulations. Both the bio-based route and the fossil-based route require a similar number and type of vessels.

3. Results and discussion

3.1. Life cycle inventory data

LCI data associated with GBL and succininc acid synthesis was compiled from commercial and open sources. The inventories for converting these intermediates to NVP were obtained through process simulations (section 2.4). In the following, these inventories are discussed to improve our understanding of the process scenarios. Initially, results are contrasted for the fossil (*Fos*) and an exemplary bio-based (*Bio-Ru-Ex*) NVP production scenario (Table 2).

While molar NVP yields are similar for the GBL-(0.65 mol_{GBL} per mol_{NVP}) and succinic acid-based (0.67 mol_{acid} per mol_{NVP}) scenario, a notable difference can be observed in substrate consumption (fossil: 1.8 kg_{substrates} per kg_{NVP} vs. bio: 2.7 kg_{substrates} per kg_{NVP}). However, the higher substrate demand in the bio-based scenario is related to four equivalents of water, formed through condensation, hydrogenation, and dehydration reactions when succinic acid is converted to NVP. On the other hand, GBL is characterized by a lower C:O ratio (higher degree of reduction), thus expelling only one equivalent of water in this value chain. While the high C:O ratio of succinic acid relates to a common challenge in making chemicals from biomass, 1 GBL itself is generated by hydrogenation of an acid anhydride and subsequent removal of co-produced water. However, the choice of our system boundaries for process simulations was not optimized towards the comparability of the utilized intermediates. These upstream steps are accounted for in the LCA.

The low amount of solvent (process water) and therefore high concentration of reactant/product streams is a clear benefit of the *Fos* scenario. Most importantly, the load on distillation columns is reduced and thus the need for steam and natural gas for heating. Yet, the conventional process scenario (*Fos*) is burdened with the expenditure of more costly utilities (electricity and sub-ambient cooling) due to the extensive use of vacuum distillations. Overall, exergy demand is thus comparable between the discussed alternatives, with 7.5% lower exergy demand for the fossil alternative.

Lastly, Fos and Bio-Ru-D differ in the amounts of generated wastewater, which are higher in the succinic acid-based production scenario. While the latter also comprises an additional inert gas purge (N₂ from dehydration), quantities of the remaining waste streams (organics and flammable gas) are comparable between Fos and Bio-Ru-D, not least due to similar overall conversion selectivity.

The scenarios for NVP production from succinic acid consider two separation strategies (distillation (-*D*) and extraction (-*Ex*)) and three catalyst systems (ruthenium-based (-*Ru*-), platinum-based (-*Pt*-) and ideal, (-*Id*-)) (Table 3). Regarding separation, oligomer removal by extraction (*Bio-Ru-Ex vs. Bio-Ru-D*) comes at the cost of an organic solvent (dichloromethane,

Table 2 Exemplary comparison of LCI data for GBL- and succinic acid-based NVP production scenarios

Fos scenario		Bio-Ru-D scenario		
Stream	Quantity	Stream	Quantity	Units
Substrates		Substrates		
γ-Butyrolactone	1.23	Succinic acid	1.71	$kg kg_{NVP}^{-1}$
Ammonia	0.26	Monoethanolamine	0.89	kg kg _{NVD} ⁻¹
Acetylene	lene 0.28 Hy		0.088	kg kg _{NVP} ⁻¹
Auxiliaries		Auxiliaries		
Process water	0.50	Process water	1.71	$kg kg_{NVP}^{-1}$
KOH	0.016	Nitrogen	0.12	$kg kg_{NVP}^{-1}$
		Ru/C	8.6×10^{4}	kg kg _{NVp} ⁻¹
		Na_2O/SiO_2	2.5×10^3	kg kg _{NVP} ⁻¹
Utilities		Utilities		
Natural gas	1.67	Natural gas	1.74	$MJ kg_{NVP}^{-1}$
Steam	2.23	Steam	9.49	$MJ kg_{NVP}^{-1}$
Cryogen ^a	1.10	Cryogen ^a	0.37	${ m MJ~kg_{NVP}}^{-1}$ ${ m MJ~kg_{NVP}}^{-1}$
Water cooling	1.40	Water cooling	1.51	$MJ kg_{NVP}^{-1}$
Air cooling	5.28	Air cooling	11.3	$MJ kg_{NVP}^{-1}$
Electricity	0.79	Electricity	0.15	kWh kg _{NVP} ⁻¹
$(Total\ exergy)^b$	(4.82)	$(Total\ exergy)^b$	(5.21)	$MJ kg_{NVP}^{-1}$
Waste ^c		Waste ^c		
		Inert gas purge	0.13	$kg kg_{NVP}^{-1}$
Flam. gas purge	0.05	Flam. gas purge	0.044	$kg kg_{NVP}^{-1}$
Organic residue	0.46	Organic residue	0.41	kg kg _{NVP} ⁻¹
Wastewater	0.78	Wastewater	2.94	kg kg _{NVP} ⁻¹

^a At T = -25 °C. ^b Calculated via the Carnot efficiency ($T_{ref} = 25$ °C) of heat at 400 °C (natural gas), 200 °C (steam), -25 °C (cryogen). ^c Exemplary stream compositions and inventories of waste treatment in the ESI (section 10†);

Table 3 Comparison of LCI data for succinic acid-based NVP production scenarios

Stream	Bio-Ru-D	Bio-Ru-Ex	Bio-Pt-Ex	Bio-Id-Ex	Unita
Stream	Quantity				Units
Substrates					
Succinic acid	1.71	1.68	1.44	1.24	kg kg _{NVP} ⁻¹
Monoethanolamine	0.89	0.87	0.74	0.64	kg kg _{NVP} ⁻¹
Hydrogen	0.088	0.086	0.057	0.047	$kg kg_{NVP}^{-1}$
Auxiliaries					
Process water	1.71	1.68	1.44	1.24	$kg kg_{NVP}^{-1}$
Dichloromethane	_	0.11	0.084	0.068	$kg kg_{NVP}^{-1}$ $kg kg_{NVP}^{-1}$
Nitrogen	0.12	0.12	0.12	0.12	kg kg _{NVP} ⁻¹
Hydrogenation catalyst ^a	8.6×10^{4}	8.4×10^{4}	7.2×10^4	6.2×10^4	kg kg _{NVP}
Na ₂ O/SiO ₂	2.5×10^3	2.5×10^3	2.5×10^3	2.5×10^3	$kg kg_{NVP}^{-1}$
Utilities					
Natural gas	1.74	1.58	1.70	3.19	MJ kg _{NVP} ⁻¹
Steam	9.49	6.93	5.68	3.25	$MJ kg_{NVP}^{-1}$ $MJ kg_{NVP}^{-1}$
Cryogen ^b	0.37	1.48	1.32	1.13	$MJ kg_{NVP}^{-1}$
Water cooling	1.51	4.62	4.01	3.55	MI kg _{NVD} ⁻¹
Air cooling	11.3	4.86	4.11	3.54	MJ kg_{NVP}^{-1}
Electricity	0.15	0.087	0.076	0.065	kWh kg _{nvp} 1
(Total exergy) ^c	(5.21)	(4.11)	(3.63)	(3.48)	$MJ kg_{NVP}^{-1}$
Waste ^d					
Inert gas purge	0.13	0.13	0.13	0.13	kg kg _{NVP} ⁻¹
Flam. gas purge	0.044	0.043	0.011	7.2×10^{3}	kg kg _{NVP} ⁻¹
Organic residue	0.41	0.22	0.18	0.11	kg kg _{NVP} ⁻¹
Wastewater	2.94	3.14	2.57	2.11	kg kg _{NVP} ⁻¹

^a Ru/C or Pt–Re/TiO₂ depending on scenario. ^b At T = -25 °C. ^c Calculated via the Carnot efficiency ($T_{\rm ref} = 25$ °C) of heat at 400 °C (natural gas), 200 °C (steam), -25 °C (cryogen). ^d Exemplary stream compositions and inventories of waste treatment in the ESI (section 10†).

DCM) and higher cooling demand due its low boiling point (~40 °C). Yet, this cost is balanced by lower steam and electricity requirements in Bio-Ru-Ex, which trace back to the heat of evaporating DCM as opposed to water and a reduced vacuum distillation need. Finally, waste quantities in the listed scenarios are affected by the purge of organic solvent and the difference in allocation of organic by-products to waste streams (see Fig. 4). For example, oligomeric by-products are allocated to organic residue and wastewater streams in distillation and extraction scenarios, respectively.

Lastly, further distinctions between Bio-x-Ex scenarios are caused by the choice of a hydrogenation catalyst, which determines overall NVP yields from succinic acid. While a recent improvement of the commercially available Ru/C catalyst, namely a bimetallic Pt-Re/TiO2 material, 41 reduces normalized process inputs by about 14% relative, further optimization (Bio-Id-Ex) would allow for 26%. For reasons of mass balancing, waste quantities are inversely affected by the implemented hydrogenation catalyst. Still, the total exergy demand of the Bio-Pt-Ex scenario is less than 5% higher than for the ideal Bio-Id-Ex scenario, indicating small room for increasing energy efficiency through catalyst development.

3.2. Life cycle assessment

The environmental impacts of succinic acid-based NVP production were determined by evaluating the LCIs described in section 3.1 with the CML2001/2016 method. Results are summarized in Fig. 6 for the global warming impact (GWI); other categories are discussed subsequently.

The global warming impact of fossil NVP production is estimated at 7.6 kg_{CO₂} eq. per kg_{NVP}. Thus, even the commercial Ru/C catalyst in the *Bio-Ru-D* scenario (5.3 kg $_{CO_2}$ eq. per kg $_{NVP}$) reduces GWI. Emission savings are mostly due to a large contribution of the fossil-derived GBL intermediate (~60%) to the global warming impact in the Fos scenario. Bio-Ru-D, on the other hand, has a larger global warming impact due to energy use (0.95 kg $_{CO_2}$ eq. per kg $_{NVP}$ vs. 0.48 kg $_{CO_2}$ eq. per kg $_{NVP}$) and waste disposal (0.83 kg_{CO}, eq. per kg_{NVP} vs. 0.37 kg_{CO}, eq. per kg_{NVP}). However, increases in utility impacts are more than compensated for by the low contribution of the succinic acid feedstock due to its biogenic origin. Avoiding four equivalents of fossil carbon per molecule of NVP lowers the global warming impact of the Bio-Ru-D scenario substantially.

Even utilizing bio-based feedstocks, the production of NVP is not carbon neutral. Remaining global warming impacts of bio-based NVP production are mainly driven by the utilization of fossil-based resources for energy provision or in the educts monoethanolamine and hydrogen. Global warming impacts could thus be decreased by utilizing renewable energy for electricity and heat generation and supplying monoethanolamine and hydrogen from renewable sources. However, such improvements would also lower the carbon intensity of the fossilbased supply chain.

Due to our cradle-to-gate system boundaries, the use phase and end-of-life of NVP is not considered. Assuming combustion with no heat recovery as the end of life of NVP would add

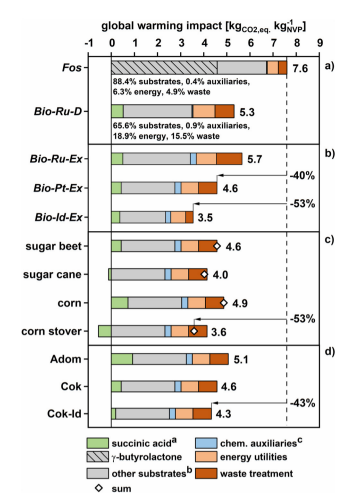


Fig. 6 Cradle-to-gate global warming impact of NVP production as a function of (a) value chain, (b) catalyst technology, (c) fermentation feedstock and (d) succinic acid production process. The process scenario is Bio-Pt-Ex, unless specified otherwise, a Succinic acid is derived from sugar beet using a process by Cok et al. 65 unless specified otherwise, $^{\rm b}$ other substrates include $\rm H_2$ and monoethanolamine (*Bio*) as well as ammonia and acetylene (Fos), catalysts (heterogeneous and KOH), solvents (H2O, DCM) and inerts (N2) are summarized as auxiliaries.

additional 2.3 kg_{CO₂} eq. per kg_{NVP} to the global warming impacts of fossil and bio-based NVP.

The overall GWI of the extractive alternative (Bio-Ru-Ex) is in-between the Bio-Ru-D case and the Fos case, with a global warming impact of 5.7 kg_{CO₂} eq. per kg_{NVP}. Compared to Bio-Ru-D, the GWI primarily increases due to the production and disposal of the extraction solvent, which is partially purged to avoid the accumulation of impurities. Yet, Bio-Ru-Ex should be considered a base case of Bio scenarios due to its conservative approach to removing oligomeric residue. Thus, changes in the catalyst system are only discussed in detail for the extraction case (Bio-x-Ex).

Here, using the more efficient Pt-Re/TiO2 catalyst as opposed to Ru/C lowers substrate consumption and waste treatment demand, thereby reducing the global warming impacts of NVP production (Bio-Pt-Ex: 4.6 kg_{CO2} eq. per kg_{NVP}). Notably, the large specific global warming impact of platinum

(19 600 kg_{CO_2} eq. per kg_{Pt}) as compared to ruthenium (3570 kg_{CO_2} eq. per kg_{Ru}) has a negligible impact due to catalyst stability and metal recycling (ESI, section $3b + 4b\dagger$). In extrapolation of this case study, a potential improvement of the Pt-catalyst (Y(HEP) = 98 mol%) would bring the global warming impact of bio-based NVP production down to 3.5 kg_{CO_2} eq. per kg_{NVP} - 53% less than in the *Fos* scenario. This change is primarily due to an efficient succinic acid conversion to NVP, which reduces normalized process input and waste output quantities. Due to these benefits, an ideal catalyst would also reduce the carbon footprint by an additional 24% compared to the best current scenario *Bio-Pt-Ex*, much more than the small improvement potential found for energy in section 3.1.

Also, fermentative succinic acid production is a rapidly developing field, with current efforts focusing, among others, on the use of alternative feedstocks. Thus, a sensitivity analysis considers four feedstocks for sugar production (Fig. 6c). While variations between cases (3.6–4.9 kg $_{\rm CO_2}$ eq. per kg $_{\rm NVP}$) are smaller than those observed for different catalyst systems, the implementation of corn stover leads to the highest GWI savings (53% relative reduction νs . Fos). This finding is expected as corn stover represents a lignocellulosic side-product of food and feed production, with limited alternative use.

Further variability in the total GWI of Bio-Pt-Ex is connected to sugar fermentation and succinic acid separation. Three scenarios for succinic acid production are compared in this regard (Fig. 6d): (i) the base case of Cok et al. 65 with glucose and energy demands of 1.35 kg and 9.9 MJ per kg of succinic acid, (ii) an alternative evaluation by Adom at al. 94 demanding 1.90 kg and 17 MJ per kg of acid and (iii) a best case, where the theoretical minimum glucose demand of 1.04 kg per kg of acid⁹⁵ is assumed within the process of Cok et al. (Cok-Id). Comparing the scenarios based on Cok et al. and Adom et al. shows a mild impact of assumptions for succinic acid production on the general advantage of Bio-Pt-Ex. Global warming impacts increase from 4.5 kg_{CO2} eq. per kg_{NVP} to 5.1 kg_{CO2} eq. per kg_{NVP} between these cases. On the other hand, pushing succinic acid fermentation to the theoretical limit could reduce the global warming impacts of the Bio-Pt-Ex scenario to $4.3 \text{ kg}_{\text{CO}_2}$ eq. per kg_{NVP}.

Finally, impacts from indirect land use change (iLUC) have recently attained increasing interest in assessing biomass-based production, especially for first-generation feedstocks. Thus, several iLUC estimates for the proposed feedstocks were derived for the succinic acid-based production scenario *Bio-Pt-Ex* considered most realistic (Fig. 7). The analysis shows that iLUC ($-0.1-0.7~{\rm kg_{CO_2}}$ eq. per kg_{NVP}) has a mild impact on the environmental merits of *Bio* scenarios, but should always be analyzed in real-life implementation.

As far as other categories are concerned, environmental impacts are estimated to decrease by up to 65% relative (*Bio-Id-Ex vs. Fos*) in the categories of photochemical ozone creation, ozone layer depletion, and abiotic depletion of fossil energy (Fig. 8). In contrast, *Bio* scenarios increase impacts in the cat-

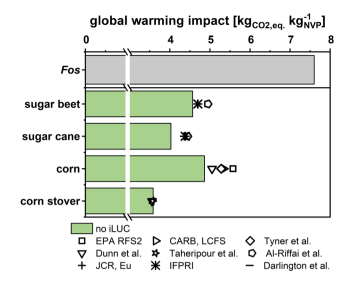


Fig. 7 Impact of different iLUC estimates on total GWI for the *Bio-Pt-Ex* scenario.^{74–81} Further information are given in the ESI section 4c.†

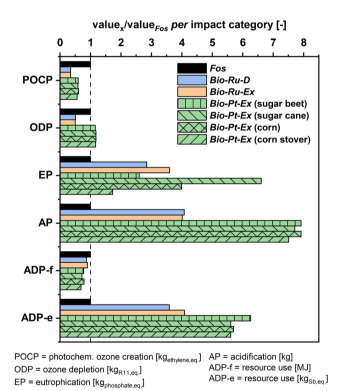


Fig. 8 Relative non-tox environmental impacts of bio-based NVP production compared to fossil-based production. Tabulated absolute values are given in the ESI section $11.\dagger$

egories of eutrophication potential, acidification potential, and abiotic resource depletion. This increase is due to biomass feedstock production in the case of eutrophication and due to the supply of noble metal catalyst components in the case of acidification and resource depletion.

Moreover, heightened eutrophication presents a commonly observed drawback in bio-based chemical production and is primarily related to direct fertilizer emissions from agriculture. Thus, surges in this category νs . Fos vary between 70 and 560% relative, depending on biomass feedstock. The minimum of this range corresponds to corn stover (US) and the maximum to sugar cane (Thailand), due to high fertilizer use in the available inventories.

In the category of acidification, increases of up to 300% for routes using ruthenium catalysts and up to 690% for routes using platinum catalysts are found. Mining and refining activities of the noble metals are the root cause and contribute between 80–90% to the total impact estimate. These contributions, in turn, mostly stem from direct sulfur dioxide emissions in the refining process, which may be avoidable through the implementation of suitable gas treatment standards. Finally, the category of abiotic resource depletion is also dominated by the catalytic material, contributing 50–80% of the total impact estimate.

This analysis highlights the need to establish a sustainable catalyst supply chain to avoid excessive acidification and to utilize waste feedstocks such as corn stover to reduce eutrophication. Yet, it must be noted that current estimates on catalyst stability are conservative due to a lack of long-term stability data. Thus, catalyst-related impacts could be lowered in the course of future technology development.

3.3. Operational cost analysis

Economic viability is another important factor in the implementation of bio-based processes. In contrast to the GWI estimates, an initial comparison of Bio-Ru-D (6.2 \$ per kg_NVP) and Fos (5.4 \$ per kg_NVP) scenarios shows higher operational costs associated with NVP from succinic acid (Fig. 9). The root cause of this difference between GWI and cost evaluations is the CO₂ benefit of biomass feedstocks, which does not directly affect their economics.

However, cost and GWI trends within the set of Bio scenarios are identical. Thus, Bio-Ru-Ex yields slightly higher costs (6.3 \$ per kg_{NVP}), mostly due to the expenditure of extraction agent and waste treatment. The cost of Bio-x-Ex can be reduced by up to 27% relative by new catalyst technologies in amidation—hydrogenation (Bio-Ru-Ex vs. Bio-Id-Ex). NVP from Bio-Id-Ex would then be produced at 15% lower cost than Fos.

The pronounced impact of catalyst selectivity on overall cost is rooted in the large contribution of substrate costs in *Fos* as well as *Bio* scenarios. For example, substrates account for 5.8 \$ per kg_{NVP} (93.5%) in *Bio-Ru-D*. This dominance of substrate cost is due to low solvent loads (see section 3.1) and straightforward separation schemes, which lead to modest energy demands. Also, waste treatment contributes more to emissions than to overall costs.

Within the substrate cost segment, succinic acid and GBL are the largest contributors, implying a strong dependence on the price of biogenic succinic acid set by a rapidly developing market. Thus, a sensitivity analysis for succinic acid price is presented in Fig. 9c (base case = 2.5 \$ per kg_{acid}). The ana-

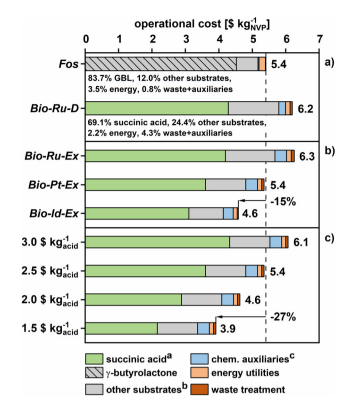


Fig. 9 Operational cost of NVP production as a function of (a) value chain, (b) catalyst technology and (c) substrate cost. 2.5 \$ per kg_{Acid} unless specified otherwise. ^a Succinic acid is derived from sugar beet, ^b other substrates include H_2 and monoethanolamine (*Bio*) as well as ammonia and acetylene (*Fos*), ^c catalysts (heterogeneous and KOH), solvents (H_2O , DCM) and inerts (N_2) are summarized as auxiliaries.

lysis shows that bio-based NVP production through the improved Pt–Re/TiO₂ catalyst (*Bio-Pt-Ex*) is likely to remain cost-competitive for succinic acid prices below 2.5 \$ per kg. This break even point equates to a price ratio of 0.69 for the utilized C4 intermediates (succinic acid/GBL, mass-based), which could be used as a rough indicator of feasibility in different GBL market conditions.

Lastly, further improvements in fermentation and separation technology have been suggested to reduce the cost of succinc acid production to 1.5 \$ per kg_{acid}. ⁸⁹ Therefore, the operational cost for *Bio-Pt-Ex* may become as low as 3.9 \$ per kg_{NVP} – almost 30% less than in the *Fos* scenario. In addition, the ESI† explores the dependence of economic viability on patent-based assumptions for fossil NVP production, showing that a fossil-based production nearing stoichiometric yield would still have higher operating costs of 4.1 \$ per kg_{NVP} (ESI, section 9†).

3.4. Uncertainty

Previous sections have discussed the environmental and economic impacts of bio-based pyrrolidone production. Therein multiple process and market scenarios, covering a broad range of potential futures were considered. Thus, the ultimate feedstock choice and availability, the separation and catalyst

technology are key variables in determining the environmental and economic performances of bio-based *N*-vinyl-2-pyrrolidone.

It is clear that the presented, scenario-based sensitivity analysis does not include all potential uncertainties for a nascent process technology. For example, the energy system in general is in a state of transition and might change drastically till an eventual implementation of bio-based pyrrolidone production. Furthermore, bio-based succinic acid production is not yet established and thus production is simulated on literature, not industrial data. On the economic side, the market development for succinic acid provides the largest uncertainty.

Despite these unkowns, the presented spread of scenarios provides a suitable error margin for the given assessment. Accordingly, there is clear potential for improved environmental impacts through the production of NVP from succinic acid, whereas the economic success is very dependent on market developments.

3.5. Relevance and future research

To the best of our knowledge, the environmental impacts of bio-based pyrrolidone production have previously remained unreported. Lammens *et al.* have investigated the economics of NVP made from glutamic acid but did not address environmental impacts. ⁹⁰ Based on simplifying assumptions, these authors found economic potential for the production of NVP and the substrate (glutamic acid) price was identified as the dominant operational cost driver. Notably, the value chain studied by Lammens and coworkers was based on decarboxylation and vinylation instead of amidation-hydrogenation and dehydration reactions. Thus, our analysis not only adds environmental aspects to the evaluation of bio-based NVP, but also expands the scope of available synthesis pathways.

The evaluation at hand can also direct future research towards an economically feasible and environmentally friendly process. In this regard, it is noted that the succinic acid conversion to NVP is a nascent technology with knowledge gaps to be addressed prior to industrial implementation. For example, the detailed properties and continuous separation behavior of oligomeric residue from amidation–hydrogenation must be explored to validate the proposed purification technologies. Moreover, DCM represents a suboptimal extraction agent in terms of performance and green chemistry, ⁹¹ which leaves potential for more benign alternatives. However, our analysis followed a conservative approach by only considering DCM due to its demonstration in the laboratory.

Our results also indicate the large potential for further catalyst research, optimizing the selectivity and long-term stability of materials for amidation-hydrogenation. Promising catalyst candidates are other Pt-MeO $_x$ combinations (*e.g.* Pt-MoO $_x$ and Pt-VO $_x$) already explored in the general context of amide reduction. ^{92,93} Due to their impact on investment cost, factors such as operation pressure and space-time-yield will also become more important as the process moves to a pilot stage.

Finally, succinic acid from fermentation must be available at a stable, competitive price to generate investment interest in succinic acid conversion technologies to bio-products. It is therefore paramount to continue research on efficient succinic acid production and separation. Methods that allow for an integration of succinic acid production and downstream conversion are also promising to lower the effective cost contribution of substrates to NVP production.

4. Conclusions

In this work, we proposed and analyzed processes for the biobased production of *N*-vinyl-2-pyrrolidone. Currently practiced fossil alternatives served as benchmark. The analysis is based on laboratory data, scientific and patent literature. Where necessary, life cycle inventories of process steps were derived from Aspen® flowsheet simulations. Finally, process scenarios were evaluated in terms of environmental impacts, such as global warming impact, and operational costs. Within these metrics, the succinic acid conversion scenario could reduce the impact/cost of NVP by up to 53% and 15%, respectively. Notably, the best current laboratory technology (*Bio-Pt-Ex*) still delivers substantial GWI reductions of 40% and reaches parity for operational cost.

Yet, economic and environmental benefits of bio-based production via succinic acid largely depend on the availability of an efficient catalyst for amidation–hydrogenation. The catalyst must convert succinic acid and ethanolamine into N-(2-hydroxyethyl)-2-pyrrolidone, while limiting the formation of side-products, such as oligomers. This challenge is currently best addressed by a tailored Pt-Re/ TiO_2 material.

On the supply side, the bio-based processes for NVP production would benefit significantly from further developments in succinic acid production from sugars and the emergence of a well-developed market for this platform chemical. Especially, lignocellulosic feedstocks would be desirable, resulting in lower succinic acid prices and reduced environmental impacts.

The study has also highlighted the potential of combining laboratory research on nascent processes with rigorous process modeling and assessment. In this regard, it should be rewarding to enter upscaling experiments for the chemical conversions and separations described herein. Respective results could be fed back, thus improving the quality of the given estimates and generating new research priorities in an iterative approach.

Author contributions

The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript. A detailed CRediT statement is provided with the submission.

Abbreviations

NVP N-Vinyl-2-pyrrolidone

GBL γ-Butyrolactone THF Tetrahydrofurane

NMP *N*-Methyl-2-pyrrolidone

2PYD 2-Pyrrolidone

Paper

LCA Life cycle assessment GWI Global warming impact

HEP N-(2-Hydroxyethyl)-2-pyrrolidone

LCI Life cycle inventory

LCIA Life cycle impact assessment
EP Eutrophication potential
AP Acidification potential
iLUC Indirect land-use change
NRTL Non-random two liquids
DCM Dichloromethane.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This work was supported by the German Federal Ministry of Education and Research (BMBF) through the projects BioPyrr and BioPVP [FKZ IBÖ-03 031B0249 and 031B0487 A] in the framework of "New Products for the Bioeconomy" (Neue Produkte für die Bioökonomie) and by funding from the European Union's Horizon 2020 research and innovation program under grant agreement no. 720695 (GreenSolRes). The authors wish to acknowledge the help of Stephanie Föller in conducting the presented experiments. Furthermore, Mr. Haus expresses his thanks towards the German Chemical Industry Fund (Fonds der chemischen Industrie, FCI), which generously supported his PhD project.

References

- 1 P. N. R. Vennestrøm, C. M. Osmundsen, C. H. Christensen and E. Taarning, *Angew. Chem., Int. Ed.*, 2011, **50**, 10502–10509.
- 2 A. Pfennig, CBEN, 2019, 6, 90-104.
- 3 Á. Galán-Martín, V. Tulus, I. Díaz, C. Pozo, J. Pérez-Ramírez and G. Guillén-Gosálbez, One Earth, 2021, 4, 565–583.
- 4 R. Meys, A. Kätelhön, M. Bachmann, B. Winter, C. Zibunas, S. Suh and A. Bardow, *Science*, 2021, 374, 71–76.
- 5 European Comission, Bioeconomy, available at: https://ec.europa.eu/info/research-and-innovation/research-area/environment/bioeconomy_en, accessed 27 September 2021.
- 6 RoadToBio Consortium, Welcome to RoadToBio, available at: https://www.roadtobio.eu/, accessed 27 September 2021.
- 7 United States Department of Agriculture, BioPreferred, available at: https://www.biopreferred.gov/BioPreferred/, accessed 27 September 2021.
- 8 T. W. House, Ind. Biotechnol., 2012, 8, 97–102.

- 9 A. Corma, S. Iborra and A. Velty, Chem. Rev., 2007, 107, 2411–2502.
- 10 A. D. Patel, K. Meesters, H. den Uil, E. de Jong, E. Worrell and M. K. Patel, *ChemSusChem*, 2013, 6, 1724–1736.
- 11 D. Kralisch, D. Ott and D. Gericke, *Green Chem.*, 2015, **17**, 123–145.
- 12 J. Becker, A. Lange, J. Fabarius and C. Wittmann, *Curr. Opin. Biotechnol.*, 2015, **36**, 168–175.
- 13 M. Montazeri, G. G. Zaimes, V. Khanna and M. J. Eckelman, ACS Sustainable Chem. Eng., 2016, 4, 6443–6454.
- 14 (a) S. Spierling, E. Knüpffer, H. Behnsen, M. Mudersbach, H. Krieg, S. Springer, S. Albrecht, C. Herrmann and H.-J. Endres, J. Cleaner Prod., 2018, 185, 476–491;
 (b) S. Y. Lee, H. U. Kim, T. U. Chae, J. S. Cho, J. W. Kim, J. H. Shin, D. I. Kim, Y.-S. Ko, W. D. Jang and Y.-S. Jang, Nat. Catal., 2019, 2, 18–33; (c) J. He, L. Chen, S. Liu, K. Song, S. Yang and A. Riisager, Green Chem., 2020, 22, 6714–6747.
- 15 E. Castro-Aguirre, F. Iñiguez-Franco, H. Samsudin, X. Fang and R. Auras, *Adv. Drug Delivery Rev.*, 2016, **107**, 333–366.
- 16 M. L. Di Lorenzo and R. Androsch, *Industrial Applications of Poly(lactic acid)*, Springer International Publishing, Cham, 1st edn., 2018, vol. 282.
- 17 S. Niphadkar, P. Bagade and S. Ahmed, *Biofuels*, 2018, **9**, 229–238.
- 18 M. Rouhany and H. Montgomery, in *Biodiesel: From Production to Combustion*, ed. M. Tabatabaei and M. Aghbashlo, Springer International Publishing, Cham, 2019, pp. 1–14.
- 19 J. J. Bozell and M. K. Patel, Feedstocks for the Future, American Chemical Society, Washington, DC, 2006, vol. 921.
- 20 T. Werpy and G. Petersen, *Top value added chemicals from biomass: volume I—results of screening for potential candidates from sugars and synthesis gas*, National Renewable Energy Lab., Golden, CO (US), 2004.
- 21 M. J. Biddy, C. Scarlata and C. Kinchin, *Chemicals from biomass: a market assessment of bioproducts with near-term potential*, National Renewable Energy Lab.(NREL), Golden, CO (United States), 2016.
- 22 J.-V. Bomtempo, F. Chaves Alves and F. de Almeida Oroski, *Faraday Discuss.*, 2017, **202**, 213–225.
- 23 S. Takkellapati, T. Li and M. A. Gonzalez, *Clean Technol. Environ. Policy*, 2018, **20**, 1615–1630.
- 24 M. Dusselier, M. Mascal and B. F. Sels, in *Selective Catalysis* for *Renewable Feedstocks and Chemicals*, ed. K. M. Nicholas, Springer, Heidelberg, 2014, pp. 1–40.
- 25 L. Corbel-Demailly, B.-K. Ly, D.-P. Minh, B. Tapin, C. Especel, F. Epron, A. Cabiac, E. Guillon, M. Besson and C. Pinel, *ChemSusChem*, 2013, 6, 2388–2395.
- 26 K. H. Kang, U. G. Hong, Y. Bang, J. H. Choi, J. K. Kim, J. K. Lee, S. J. Han and I. K. Song, *Appl. Catal.*, A, 2015, 490, 153–162.
- 27 C. S. Spanjers, D. K. Schneiderman, J. Z. Wang, J. Wang, M. A. Hillmyer, K. Zhang and P. J. Dauenhauer, *ChemCatChem*, 2016, 8, 3031–3035.

- 28 X. Di, C. Li, B. Zhang, J. Qi, W. Li, D. Su and C. Liang, *Ind. Eng. Chem. Res.*, 2017, **56**, 4672–4683.
- 29 X. Di, C. Li, G. Lafaye, C. Especel, F. Epron and C. Liang, *Catal. Sci. Technol.*, 2017, 7, 5212–5223.
- 30 A. Cukalovic and C. V. Stevens, *Biofuels, Bioprod. Biorefin.*, 2008, 2, 505–529.
- 31 weastra s.r.o. WP 8.1. Determination of market potential for selected platform chemicals, available at: https://www.cbp.fraunhofer.de/content/dam/cbp/en/documents/project-reports/BioConSepT_Market-potential-for-selected-platform-chemicals_report1.pdf, accessed 13 October 2021.
- 32 J.-P. Lange and S. H. Wadman, *ChemSusChem*, 2020, **13**, 5329–5337.
- 33 M. J. Hülsey, H. Yang and N. Yan, *ACS Sustainable Chem. Eng.*, 2018, **6**, 5694–5707.
- 34 J. F. White, J. E. Holladay, A. A. Zacher, J. G. Frye and T. A. Werpy, *Top. Catal.*, 2014, 57, 1325–1334.
- 35 G. Budroni and A. Corma, J. Catal., 2008, 257, 403-408.
- 36 W. Fischer, D. Klein, A. Künkel, R. Pinkos and E. Scholten, *US Patent*, 8017790B2, 2011.
- 37 T. Thosukhowong, K. Roffi, S. R. More, R. L. Gustine and S. Tanielyan, *WO Patent*, 2013033649A1, 2013.
- 38 brgfx, Vector Image Tree. Hintergrund Vektor erstellt von brgfx de.freepik.com , available at: https://de.freepik.com/brgfx, accessed 17 October 2021.
- 39 M. O. Haus, Y. Louven and R. Palkovits, *Green Chem.*, 2019, 21, 6268–6276.
- 40 H. Krishna, M. O. Haus and R. Palkovits, *Appl. Catal., B*, 2021, 286, 119933.
- 41 M. O. Haus, A. Meledin, S. Leiting, Y. Louven, N. C. Roubicek, S. Moos, C. Weidenthaler, T. E. Weirich and R. Palkovits, *ACS Catal.*, 2021, **11**, 5119–5134.
- 42 (a) International Organization for Standardization, ISO 14040:2006, available at: https://www.iso.org/standard/37456.html, accessed 29 September 2021; (b) L. Ismail, J. Alcorta and N. Perou, *The computational structure of life cycle assessmnet. Dordrecht*, Kluwer (Eco-efficiency in industry and science, v. 11), London, 2002.
- 43 M. Weiss, J. Haufe, M. Carus, M. Brandão, S. Bringezu, B. Hermann and M. K. Patel, *J. Ind. Ecol.*, 2012, **16**, 169–181.
- 44 O. G. Griffiths, R. E. Owen, J. P. O'Byrne, D. Mattia, M. D. Jones and M. C. McManus, *RSC Adv.*, 2013, 3, 12244– 12254.
- 45 I. Muñoz, K. Flury, N. Jungbluth, G. Rigarlsford, L. M. i Canals and H. King, *Int. J. Life Cycle Assess.*, 2014, 19, 109–119.
- 46 H. de Bruijn, R. van Duin, M. A. J. Huijbregts, J. B. Guinee, M. Gorree, R. Heijungs, G. Huppes, R. Kleijn, A. de Koning, L. van Oers, A. Wegener Sleeswijk, S. Suh and H. A. Udo de Haes, *Handbook on Life Cycle Assessment*, Springer Netherlands, Dordrecht, 2002, vol. 7.
- 47 N. von der Assen, J. von Jung and A. Bardow, *Energy Environ. Sci.*, 2013, **6**, 2721.
- 48 N. von der Assen, P. Voll, M. Peters and A. Bardow, *Chem. Soc. Rev.*, 2014, 43, 7982–7994.

- 49 R. Pelton, Int. J. Life Cycle Assess., 2019, 24, 12-25.
- 50 B. Winter, R. Meys and A. Bardow, J. Cleaner Prod., 2021, 290, 125818.
- 51 C. Andersson, D. Hodge, K. A. Berglund and U. Rova, *Biotechnol. Prog.*, 2007, 23, 381–388.
- 52 S. Chan, S. Kanchanatawee and K. Jantama, *Bioresour. Technol.*, 2012, **103**, 329–336.
- 53 C. Li, K. L. Ong, Z. Cui, Z. Sang, X. Li, R. D. Patria, Q. Qi, P. Fickers, J. Yan and C. S. K. Lin, *J. Hazard. Mater.*, 2021, 401, 123414.
- 54 W. Schwarz, J. Schossig, R. Rossbacher and H. Höke, *in Ullmann's Encyclopedia of Industrial Chemistry*, Wiley-VCH Verlag GmbH & Co. KGaA, Weinheim, Germany, 2000.
- 55 A. L. Harreus, R. Backes, J.-O. Eichler, R. Feuerhake, C. Jäkel, U. Mahn, R. Pinkos and R. Vogelsang, in Ullmann's Encyclopedia of Industrial Chemistry, Wiley-VCH Verlag GmbH & Co. KGaA, Weinheim, Germany, 2000.
- 56 M. Rudloff, P. Stops, E. Henkes, H. Schmidtke, R.-H. Fischer, M. Julius, R. Lebkücher and K.-H. Ross, US Patent, 7164031B2, 2007.
- 57 W. Staffel, S. Käshammer, R. Kessinger, R. Vogelsang, A. Paul and L. Tuttelberg, *WO Patent*, 2009071479A1, 2009.
- 58 R. Vogelsang, S. Käshammer, W. Staffel, U. Eiden, A. Brand and L. Tuttelberg, *WO Patent*, 2009074534A1, 2009.
- 59 K.-C. Liu and P. D. Taylor, US Patent, 4824967A, 1989.
- 60 K.-C. Liu and P. D. Taylor, US Patent, 4873336A, 1989.
- 61 B. Puetzer, L. Katz and L. Horwitz, US Patent, 2775599A, 1956.
- 62 K. Ebel, M. Eiermann, T. Narbeshuber and E. Gehrer, *US Patent*, 5994562A, 1999.
- 63 Y. Shimasaki, H. Yano, H. Sugiura and H. Kambe, BCSJ, 2008, 81, 449–459.
- 64 X. Li, W. Wan, J. G. Chen and T. Wang, ACS Sustainable Chem. Eng., 2018, **6**, 16039–16046.
- 65 B. Cok, I. Tsiropoulos, A. L. Roes and M. K. Patel, *Biofuels, Bioprod. Biorefin.*, 2014, **8**, 16–29.
- 66 Sphera, GaBi Software, available at: https://gabi.sphera.com/deutsch/software/gabi-software/, accessed 10.16.2021.
- 67 IHS Markit, Chemical Economics Handbook, 2017.
- 68 C. Seyler, T. B. Hofstetter and K. Hungerbühler, *J. Cleaner Prod.*, 2005, **13**, 1211–1224.
- 69 A. Köhler, PhD Thesis, ETH Zurich, 2006. Available from DOI: 10.3929/ethz-a-005206273, last visited 11.05.2022.
- 70 P. Pawelzik, M. Carus, J. Hotchkiss, R. Narayan, S. Selke, M. Wellisch, M. Weiss, B. Wicke and M. K. Patel, *Resour.*, *Conserv. Recycl.*, 2013, 73, 211–228.
- 71 S. E. Tanzer and A. Ramírez, *Energy Environ. Sci.*, 2019, **12**, 1210–1218.
- 72 T. Searchinger, R. Heimlich, R. A. Houghton, F. Dong, A. Elobeid, J. Fabiosa, S. Tokgoz, D. Hayes and T.-H. Yu, *Science*, 2008, **319**, 1238–1240.
- 73 J. A. Mathews and H. Tan, *Biofuels, Bioprod. Biorefin.*, 2009, 3, 305–317.
- 74 W. E. Tyner, F. Taheripour, Q. Zhuang, D. Birur and U. Baldos, Land Use Changes and Consequent CO2 Emissions due to US Corn Ethanol, Purdue University, 2010.

75 L. Marelli, F. Ramos, R. Hiederer and R. Koeble, Estimate of GHG emissions from global land use change scenarios, 2011.

- 76 D. Laborde Debucquet, Assessing the land use change consequences of European biofuel policies, 2011.
- 77 California Air Resources Board, LCFS Land Use Change Assessment, available at: https://ww2.arb.ca.gov/resources/ documents/lcfs-land-use-change-assessment, accessed 17 October 2021.
- 78 P. Al-Riffai, B. V. Dimaranan and D. Laborde Debucquet, Global trade and environmental impact study of the EU biofuels mandate, 2010.
- 79 T. Darlington, D. Kahlbaum, D. O'Connor and S. Mueller, Land Use Change Greenhouse Gas Emissions of European Biofuel Policies Utilizing the Global Trade Analysis Project (GTAP) Model, 2013.
- 80 F. Taheripour and W. E. Tyner, *Induced Land Use Emissions due to First and Second Generation Biofuels and Uncertainty in Land Use Emission Factors*, Economics Research International, 2013.
- 81 J. B. Dunn, S. Mueller, H.-Y. Kwon and M. Q. Wang, *Biotechnol. Biofuels*, 2013, 6, 51.
- 82 macrovector, Vector Image Industrial Building Icons. Abstrakt
 Vektor erstellt von macrovector de.freepik.com , available at: https://de.freepik.com/macrovector, accessed 17
 October 2021.
- 83 Y. Yamaguchi, H. Yano, K. Hitoshi, A. Kurusu and Y. Shimasaki, *EP Patent*, 1094061B2, 2001.
- 84 S. W. Benson, Thermochemical kinetics, Wiley, 1976.

- 85 A. Klamt, F. Eckert and W. Arlt, *Annu. Rev. Chem. Biomol. Eng.*, 2010, 1, 101–122.
- 86 ChemCatBio, CatCost, available at: https://catcost.chemcatbio.org/, accessed 13 October 2021.
- 87 E. Mancini, S. S. Mansouri, K. V. Gernaey, J. Luo and M. Pinelo, Crit. Rev. Environ. Sci. Technol., 2020, 50, 1829– 1873.
- 88 A. Broch, S. K. Hoekman and S. Unnasch, *Environ. Sci. Policy*, 2013, **29**, 147–157.
- 89 M. Morales, M. Ataman, S. Badr, S. Linster, I. Kourlimpinis, S. Papadokonstantakis, V. Hatzimanikatis and K. Hungerbühler, *Energy Environ. Sci.*, 2016, 9, 2794– 2805.
- 90 T. M. Lammens, S. Gangarapu, M. C. Franssen, E. L. Scott and J. P. Sanders, *Biofuels, Bioprod. Biorefin.*, 2012, 6, 177– 187.
- 91 F. P. Byrne, S. Jin, G. Paggiola, T. H. M. Petchey, J. H. Clark, T. J. Farmer, A. J. Hunt, C. Robert McElroy and J. Sherwood, SustainableChem. Processes, 2016, 4, 7.
- 92 K. Shimizu, W. Onodera, A. S. Touchy, S. M. A. H. Siddiki, T. Toyao and K. Kon, *ChemistrySelect*, 2016, 1, 736–740.
- 93 T. Mitsudome, K. Miyagawa, Z. Maeno, T. Mizugaki, K. Jitsukawa, J. Yamasaki, Y. Kitagawa and K. Kaneda, *Angew. Chem., Int. Ed.*, 2017, **56**, 9381–9385.
- 94 F. Adom, J. B. Dunn, J. Han and N. Sather, *Environ. Sci. Technol.*, 2014, **48**, 14624–14631.
- 95 T. Tiso, T. Narancic, R. Wei, E. Pollet, N. Beagan, K. Schröder, A. Honak, M. Jiang, S. T. Kenny, N. Wierckx, R. Perrin, L. Avérous, W. Zimmermann, K. O'Connor and L. M. Blank, *Metab. Eng.*, 2021, 66, 167–178.