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Heterogeneous catalytic conversion of solid anaerobic digestate waste to biofuels and value-added chemicals†

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Increasing use of anaerobic digestion (AD) in agri-food waste management for the production of renewable methane has presented issues surrounding AD. One of the challenges of AD is handling the solid fraction of the digestate. Whilst the digestate, especially the liquid fraction, can currently be used as bio-fertiliser, there are still limitations to its use because of regulations on the nutrient loading in soil and possible run off resulting in eutrophication of water. Hence, there is a need for alternative forms of valorisation of both the liquid and solid digestate (SD). This work will focus on the potential of SD being a feedstock for biofuel and fine chemicals. The valorisation of the lignin constituent of SD would be a significant contribution to the success of the lignocellulosic biorefinery economy. This paper investigates the hydrogenolysis (HDO) of SD using Ni/ZnO₂, Cu/Al₂O₃ and platinum group metal (PGM) catalysts such as Rh/C, Pd/C and Ru/C to extract the biofuel. The major products from the depolymerisation of the phenolic lignin monomer constituent of the SD are the known octane booster ethylguaiacol and the structurally related antioxidant butylated hydroxytoluene. Furthermore, using 2-methyltetrahydrofuran as a green solvent increased substrate conversion (by ~13%) and bio-oil yield (by ~17 wt%) compared to dioxane solvent used in previous studies. The life cycle assessment of the catalytic extraction of anaerobic digestate showed a total energy consumption of 849 MJ per tonne of digestate utilised and a net energy ratio of 1.7, which is considerably better than fossil petrol and diesel and comparable to conventional temperate biofuels.

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Introduction

The global energy demand is still largely met with fossil-derived fuels.^{1–3} Furthermore, many everyday use chemicals are obtained from petroleum which is a non-renewable natural resource.^{4,5} Biomass offers a potential sustainable route for meeting future energy and chemical demands^{6,7} as a renewable alternative. Unlike other renewable synthesis routes, biomass has the

potential to provide both aromatic and aliphatic compounds to produce biofuels, chemicals and polymers without numerous processing steps.^{8,9} Biomass feedstock known as lignocellulosic biomass comprises three main components: cellulose, hemicellulose and lignin at 40:30:30 wt%, respectively (on average).¹⁰ While cellulosic materials are broken down in AD, lignin-based materials tend not to be, due to their physical structure, which has a highly disorganised and chemically varied structure bonded by aryl ether, β -O-4 and α -O-4 linkages. As such, residual digestate can have high lignin content.¹¹ The targeting of lignin as a sustainable source of aromatic intermediates has been a major goal of biomass valorisation for several decades, and recent rational approaches have shown some success.¹² Furthermore, there has been progress in integrating fuels and chemicals into a wider biorefinery model,^{13,14} which opens options for all levels of added value as part of a biorefinery mix.

Digestate is currently used in the agricultural sector to condition and improve the soil; however, with supplies typically exceeding local demands, the surplus digestate needs to be transported to areas lacking in nutrients or disposed of using

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additional downstream processing. Studies have shown that the transportation costs to the location of usage exceeds its value of being used as fertiliser^{15,16} and adds to its carbon footprint. Also, high nitrogen content, pathogens and chemical oxygen demand (COD) generated by digestate could be troublesome when leached into groundwater. Hence, the control of digestate needs to be mandated due to the pollution and disposal challenges it can cause.^{17–19}

Residual digestate contains significant quantities of water, which means that the digestate is most often divided into two distinct fractions (liquid and wet solid) before further usage.^{20,21} Multiple studies have shown improved techniques for digestate application. Mangwandi *et al.* and Rehl *et al.*^{22,23} reported the use of digestate to produce granular organic fertiliser and compost, while Cathcart *et al.* investigated the use of the digestate solid fraction as a fuel pellet.²⁴ Waste streams from AD could also be valorised as a nutrient for microalgae to produce a protein alternative to soybeans.²⁵

Akor *et al.* investigated the potential of a circular economy at a 500 kW standalone on-farm AD plant that produces 102,429 MJ per day of biogas from a combination of three feedstocks (grass silage, chicken litter and cattle slurry at 20.5, 8.5 and 13.7 tonnes per day, respectively).²⁶ This biogas production was found to generate ~60 kg of solid digestate per tonne of feedstock.²⁷ As such, based on the lower heating value, this would indicate the potential of pyrolysis of SD to produce an additional 653.4 kJ per tonne of feedstock, meaning a potential increase of at least 27.9 MJ energy production per day.²⁶ Post pyrolytic char is expected to have become even more concentrated in inorganic metals, such as potassium, indicating that this char can become a viable feedstock for the fertiliser industry, as has been previously reported.²⁸ This understanding aids the concept of a circular economy for anaerobic digestion and the energy potential of SD, thereby making AD plants more profitable.

During the last decade, the conversion of polysaccharides into renewable chemicals has received much attention,^{29,30} but the technologies for lignin valorisation are still less well-developed despite the progress already made. This is attributed to the intrinsic difficulty and heterogeneity of the aromatic biopolymer.³¹ Lignin is a highly branched polymer interlinked with di-phenyl ether units consisting of the building blocks *p*-coumaryl, coniferyl and sinapyl alcohol.^{32,33} The predominant bond connecting the building blocks is the β -O-4 linkage, next to C–C bonds.³⁰ As the biggest renewable origin of phenolic biopolymer with distinctive chemical structure, lignin is a promising feedstock for bulk and fine chemicals as well as fuels.^{31,32,34} Its uses are as dispersants, emulsifiers, and ingredients in resins and plastic products^{31,35} as well as the production of fine chemicals, such as vanillin.^{36,37}

The production of platform chemicals from the depolymerisation of biomass, especially lignin, is still a big challenge.^{31,34,35} One promising route in depolymerising lignin to produce low-molecular-weight compounds is *via* selective catalytic processes.³¹ Despite the complexity involved in catalytic depolymerisation of lignin, initial results are encouraging.^{38–40} Methods considered as

significant depolymerisation processes are hydrolysis, hydrogenolysis, gasification, pyrolysis, liquid-phase reforming, and chemical oxidation.^{26,32,41,42} These methods normally yield modest amounts of various monomer compounds. Cho *et al.*⁴³ reported the use of catalytic or noncatalytic fast pyrolysis and base-catalysed lignin depolymerisation (BCD) complex bio-oils with good quantity of 4-alkylated and methoxylated phenols, as well as a small yield of syringols, guaiacols, and catechols.⁴⁴ Reductive depolymerisation such as hydrogenolysis and liquid phase reforming is another promising route for lignin valorisation.⁴⁵ Different authors have reported high yields of phenolic monomers, up to 55%.^{46–49} Garrett *et al.*⁵⁰ used hydrogenolysis to depolymerise suberin from bark which is the biopolymer source that is hydrophobic in nature. Garrett *et al.*⁵⁰ also reported the depolymerisation of softwood using a heterogeneous metal catalyst such as Rh/C and Pd/C to produce 4-propylguaiacol and dihydroconiferyl alcohol bio-aromatic monomers. McCallum *et al.*⁵¹ studied the effect of base and a different green solvent such as methyl tetrahydrofuran (MeTHF) to Rh/C catalysed depolymerisation of suberin and the resultant product yield. Zakzeski *et al.*³² investigated and reported the catalytic production of monomeric and dimeric molecules from softwood. Schutyser *et al.*⁵² reported the use of selective nickel-catalysed (Ni/ZrO₂, Ni/CeO₂, Ni/Al₂O₃), Ru/C and Cu/ZrO₂ hydrogenolysis to convert lignin-derived phenolic compounds to cyclohexanone-based polymer building blocks. It also said that nickel-based catalysts are cheaper compared to noble metals (*e.g.* Rh, Pd, Ru, Au, *etc.*).⁵² Also, Ni, Cu and Fe are more green and sustainable alternatives compared to the heavy metals; however, the so-called benign metals (Ni, Cu *etc.*) could be toxic as well, depending on their chemical reactivity.^{53–55} Mortensen *et al.*⁵⁶ investigated and reported the use of Cu/Al₂O₃, Ni/ZrO₂, Pd/C, Pt/C, and Ni–V₂O₅/SiO₂ for HDO of phenol as a model compound for bio-oil. Roberts *et al.*⁴⁰ examined and reported modest yields of bio-phenolics such as syringols, guaiacols and catechols in base-catalysed depolymerisation (BCD) of lignin.

Herein, the potential of biofuel production from the catalytic depolymerisation of SD from an AD plant which contains 29% lignin (as determined by the Klason analysis of insoluble acid lignin; see Experimental section below) in the lignocellulosic material was investigated using a range of heterogeneous precious and active metal catalysts in an autoclave batch reactor. Additionally, this paper also uses life cycle assessment (LCA) to understand energy balance of the conversion of 1 tonne of anaerobic digestate to biofuels.

Experimental

Solid digestate preparation and characterisation

Digestate samples were collected from AgriAD and Agri-Food and Biosciences Institute (AFBI) AD plants located in Northern Ireland. Each digestate sample (1 L) was separated into liquid and solid using a centrifuge B 4i model operated at 6000 rpm for 9 minutes. The liquid and solid fractions were both then stored in the fridge at 0 °C overnight before being freeze-dried



(Lyotrap freeze dryer operated at $-50\text{ }^{\circ}\text{C}$ and 0.1 mbar vacuum pressure) to completely evaporate the moisture. The freeze-dried samples were ground to a 1 mm particle size using a Kinematica Polymix grinder (PX-MFC 90D) for the hydrogenolysis reaction.

Prior to reaction, Klason Insoluble Acid (KIL)⁵⁵ analysis was carried out with the sample to confirm the percentage of lignin and was calculated to be $\sim 29\%$. KIL analysis was run in duplicate (calculation shown in ESI†).

2-Methyltetrahydrofuran (2-MeTHF) and aluminium chloride (AlCl_3 , 99%) were supplied by Alfa Aesar, while ammonia solution (35%) and copper oxalate hemihydrate ($\text{CuC}_2\text{O}_4 \cdot 1/2\text{H}_2\text{O}$, 98%) were supplied by Sigma Aldrich UK. All reactions were done using doubly distilled water. Commercial rhodium, palladium and ruthenium on carbon catalysts (5 wt% Rh 64%wet, 5 wt% Pd, Ru 60%wet) were supplied by Johnson Matthey. All other reagents and analytical standards used for GC-MS derivatisation were purchased from Sigma Aldrich.

Catalyst preparation

Alumina support preparation. The preparation of the alumina support has been described elsewhere.⁵⁷ It was prepared from aluminium chloride anhydride that was then precipitated by ammonia solution; the resulting precipitate was calcined at $550\text{ }^{\circ}\text{C}$ for 4 h, at a heating rate of $2\text{ }^{\circ}\text{C min}^{-1}$. This was heated in ceramic crucibles in a muffle oven (Carbolite ELF 11/6 laboratory chamber furnace, 6 L capacity).

Copper catalysts preparation. Metal-loaded catalysts were prepared by incipient wet impregnation with the aid of sonication for effective mixing. To prepare 6 wt% $\text{Cu}/\text{Al}_2\text{O}_3$, copper oxalate hemihydrate (10 g) was dissolved with 0.5 g of support in $\sim 5\text{ mL}$ of deionised water. This was sonicated at $80\text{ }^{\circ}\text{C}$ (Crest ultrasonic bath model 200 HT) at a frequency of 45 kHz, leading to a homogeneous phase. All mixtures were then evaporated to dryness. The 6% $\text{Cu}/\gamma\text{-Al}_2\text{O}_3$ catalyst was then calcined at $550\text{ }^{\circ}\text{C}$.

Nickel catalysts preparation. The nickel-supported catalysts were prepared as described elsewhere⁵² through incipient wetness impregnation of the oxide with an aqueous solution of $\text{Ni}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$ and dried at $60\text{ }^{\circ}\text{C}$ for 24 h. The samples were reduced at $550\text{ }^{\circ}\text{C}$ (heating rate $5\text{ }^{\circ}\text{C min}^{-1}$) for 1 h under a flow of H_2 ($120\text{ mL min}^{-1}\text{ g}^{-1}$).

Catalyst characterisation

Powdered X-ray diffraction (XRD) was performed on used catalysts using a PANalytical X'pert Pro X-ray diffractometer. The diffractometer was fitted with $\text{CuK}\alpha$ X-ray source with a wavelength of $1.5405\text{ \AA}$. Diffractograms were collected from 20° to 80° with a step size of 0.01° . The X-ray tube was operated at 40 kV and 40 mA. Selected peaks were characterised by comparing with diffraction patterns using the automatic search function reference database in the X'pert software library.⁵⁵

Brunauer-Emmett-Teller (BET) analysis (using Micromeritics ASAP 2020 system) was carried out to measure the surface area and pore volume of the used catalysts. Measurement was done by N_2 adsorption. The was cleaned using nitrogen flow,

heated initially to $90\text{ }^{\circ}\text{C}$ at a rate of $15\text{ }^{\circ}\text{C min}^{-1}$ and held at this temperature for 1 h; before being heated to $125\text{ }^{\circ}\text{C}$ at the same ramp rate and held at this temperature for 28 h.

Hydrolysis reaction

Hydrogenolysis of SD was conducted under the same batch conditions reported in the previous study.⁴⁸ To summarise, the catalysts (Rh/C Pd/C Ru/C, $\text{Cu}/\text{Al}_2\text{O}_3$, Ni/ZrO_2 ; 0.5 g), SD powder (0.1 mm, 3.0 g) and 44 mL of the solvent (2-MeTHF and distilled water, in a ratio of 1:1), were added together in a reactor (Hastelloy[®] 100 mL volume, Autoclave Engineers). The reactor chamber was purged with hydrogen prior to the reaction. In the reaction, the pressure was set to 40 bar H_2 with the stirring rate set to 1000 rpm. The initial temperature was set at $100\text{ }^{\circ}\text{C}$ and thereafter increased at a rate of $10\text{ }^{\circ}\text{C min}^{-1}$ to $200\text{ }^{\circ}\text{C}$ and then dwelled for 1 h.

Product extraction and characterisation

Product extraction methods. The reactor was cooled to room temperature, and the resultant reaction mixture was filtered under vacuum. The filter cake was washed with 1:1 2-MeTHF: water ($3 \times 20\text{ mL}$). The combined filtrates were extracted with 2-MeTHF ($3 \times 100\text{ mL}$), dried over anhydrous magnesium sulphate, and concentrated under vacuum resulting in the isolation of the bio-oil. This bio-oil was dissolved in 1:1 2-MeTHF: water (30 mL) to which sodium hydroxide (0.5 g) was added to neutralise free acids. After refluxing for 4 h, the biphasic mixture was extracted with 2-MeTHF ($2 \times 100\text{ mL}$).

Fourier transform infra-red (FTIR) analysis. The functional groups present in the extracted bio-oil were determined using Agilent Technologies Cary 630 FTIR attached with attenuated total reflectance (ATR). For each spectrum, 5 mg of the sample was analysed. For each IR spectra, 40 scans were collected and averaged (step size of 10 cm^{-1} in the range of $600\text{--}4000\text{ cm}^{-1}$).

Gas chromatography-mass spectrometry (GC-MS) analysis. The bio-oil composition was analysed using Agilent Technologies 7890B – 5977A GC – MSD with PerkinElmer Turbomass detector and BP5 column. Sample preparation required that 20 mg of extracted bio-oil was dissolved in a 2 mL solution of (1.25 M) hydrochloric acid (HCl) in anhydrous methanol. The mixture was heated under reflux for 4 h under nitrogen gas, after which the methanol was removed by evaporation using a nitrogen stream. The residue was suspended in water (5 mL) and extracted with 2Me-THF ($3 \times 5\text{ mL}$). Using a nitrogen stream, the organic layer was dried and concentrated to an oily residue. This was further dried under vacuum for 2 h. After this, the bio-oil was dissolved in 0.3 mL anhydrous pyridine, and 0.5 mL *N*, *O*-bis(trimethylsilyl)trifluoroacetamide (containing 10% trimethylsilane) was then added. This mixture was heated at $70\text{ }^{\circ}\text{C}$ with agitation for 1 h under nitrogen. After cooling, 0.2 mL of 1 mg mL^{-1} (trimethylsilyl)cholesterol was added as an internal standard. This derivatised mixture (1.2 mL) was injected directly into the GC-MS for characterisation.

Nuclear magnetic resonance (NMR) analysis. The same protocol from previous studies⁵⁰ using ^1H NMR spectroscopy was quantitatively carried out on the extracted bio-oil (Bruker



Avance spectrometer at 300 MHz) with tetramethylsilane (TMS) or vanillin as internal standard, as specified below. 200 mg of the extracted bio-oil was dissolved in a mixture of 10 mL of deuteriated chloroform and 0.2 mL of deuteriated methanol. Vanillin (1 mg mL^{-1}) in deuteriated chloroform was used as an internal standard. Dissolved bio-oil was mixed 1:1 with internal standard (total 2 mL) and measured into a sample vial for NMR analysis. Quantification of the extracted bio-oil was done by comparing the bio-oil peaks with the aldehydic internal standard.

Life cycle assessment: anaerobic digestate conversion to biofuels

LCA is a tool used to quantify system-wide environmental burdens of products, processes and technologies.⁵⁸ The LCA framework and methodology applied in the present study is in accordance with ISO 14040 guidelines (ISO, 2006) and recommendations presented by Reference Life Cycle Data System (ILCD) Handbook.⁵⁹ In this study, the goal of using LCA was to quantify the energy balance of the conversion of anaerobic digestate to biofuels. The functional unit was considered as 1 tonne of anaerobic digestate. The digestate treatment options were tracked from digestate origin at the biogas plant *via* the treatment process. LCA was conducted using a cradle-to-gate attributional approach and excluded environmental impacts due to infrastructure processes. The following life-cycle stages were considered: (1) transportation of anaerobic digestate to treatment plant; (2) digestate dewatering and drying; (3) conversion of solid digestate (from Stage 2) to biofuels *via* catalytic hydrogenolysis (Fig. 1). Production of anaerobic digestate was not included in this system, as it is a waste-based resource and present in abundance with anaerobic digestion plants.

Results and discussion

Catalyst characterisation

Surface area measurements. The surface area measurements from the BET analysis are shown in Table 1. The surface areas

Table 1 Brunauer–Emmett–Teller (BET) physical characteristics of catalysts used in the reaction

Catalysts	Surface area ($\text{m}^2 \text{g}^{-1}$)	Pore volume ($\text{cm}^3 \text{g}^{-1}$)	Particle size (nm)
Rh/C	809	0.66	19.9
Pd/C	1179	1.04	14.3
Ni/ZrO ₂	345	0.08	4.8
Cu/Al ₂ O ₃	257	0.33	4.6

of Rh/C, Pd/C, Ni/ZrO₂ and Cu/Al₂O₃ were 809, 1179, 345 and $257 \text{ m}^2 \text{g}^{-1}$, respectively. The two catalysts, Ni/ZrO₂ and Cu/Al₂O₃, that were prepared by incipient wet impregnation were found to have a lower surface area of 345 and $257 \text{ m}^2 \text{g}^{-1}$ than the PMG catalysts, Rh/C and Pd/C, which were 809 and $1179 \text{ m}^2 \text{g}^{-1}$, respectively. As expected, the pore volume of Rh/C and Pd/C (0.66 and $1.04 \text{ cm}^3 \text{g}^{-1}$, respectively) was greater than Ni/ZrO₂ and Cu/Al₂O₃ (0.08 and $0.33 \text{ cm}^3 \text{g}^{-1}$, respectively). The trend in metal particles within the catalysis is Rh/C > Pd/C > Ni/ZrO₂ > Cu/Al₂O₃ with a particle size of 19.9, 14.3, 4.8 and 4.6 nm, respectively.

XRD analysis. The diffractograms of the used catalysts are shown in Fig. 2. For the Cu/Al₂O₃ catalysts, the presence of alumina support structure was observed by the diffraction peaks at $2\theta = 35.0, 38.0, 45.0$ and 66.0° (JCD no. 04-0875).⁶⁰ For the Ni/ZrO₂, the peaks at 2θ equals $29.5, 33.0, 49.0$ and 58.5° were assigned to ZrO₂ (JCPDS no. 38-1436),^{60,61} and observed diffraction peaks at 2θ equal $37.0, 44.0$, and 62.4° were assigned to NiO.⁶² Peaks of 2θ at $40.0, 45.0$ and 68.0° corresponded to metallic Pd⁶³ in the Pd/C catalysts, while the broad peak at 40° of the Rh/C diffractogram was assigned to Rh (JCPDS no. 05-0685).⁶⁴

FTIR analysis of the bio-oil

The FTIR spectra of the SD and Biochar (Johnson Matthey) are shown in Fig. 3. While lignin structures and potential depolymerisation products can vary considerably, there are a number of similarities and also some differences between the bio-oil produced in this study compared with previously reported results.

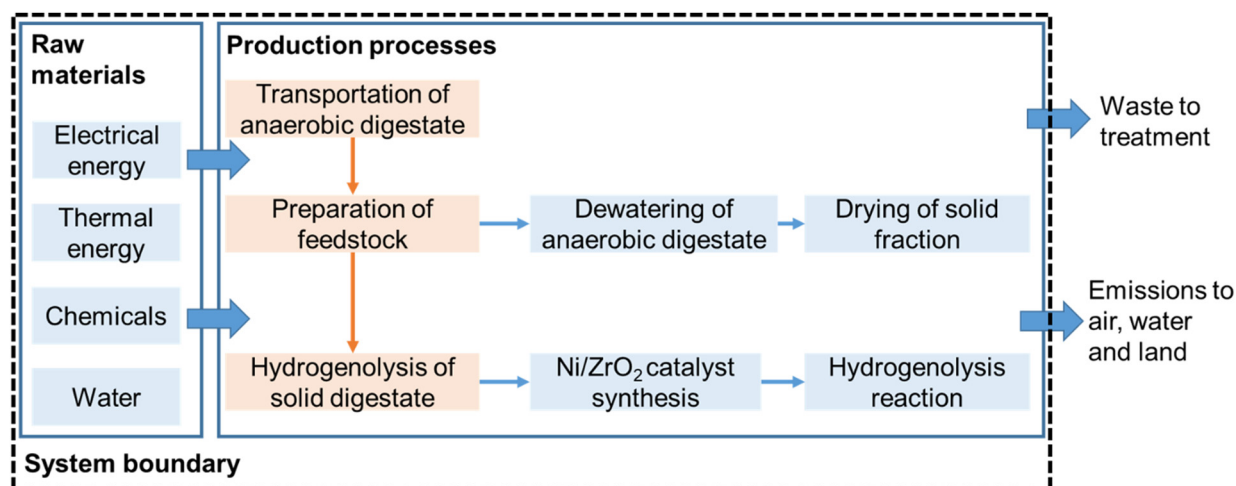


Fig. 1 System boundary considered for computing life cycle environmental impacts of the conversion of 1 tonne of digestate.



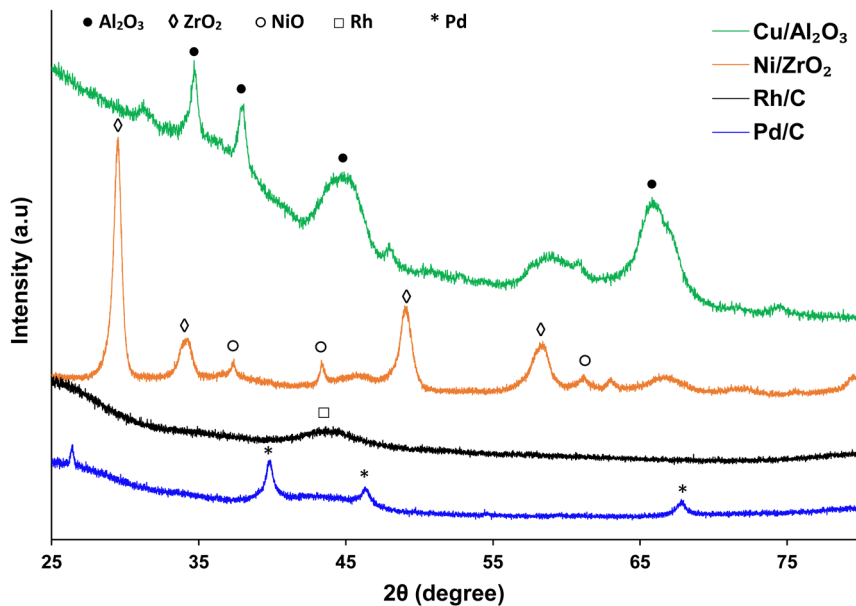


Fig. 2 XRD plots of the catalysts used within the study.

The absorption bands at 1250 and 3300 cm^{-1} are ascribed to the C–C and C–H stretching of cellulose and lignin, and carboxylic and phenolic functional group, respectively.⁶⁵ The presence of O–H vibrations between 3050 and 3600 cm^{-1} , and C=O stretching vibration shown between 1650 and 1850 cm^{-1} , confirms that carboxylic acids and their derivatives are present in the spectrum as would be expected from the literature. Pakdal and Roy^{65,66} reported a number of carboxylic acids (formic and acetic acids) detected in oils from the pyrolysis of wood. The absorbance between 1650 – 1850 cm^{-1} with C=O stretching vibrations also reveals that ketones and aldehydes are present.⁶⁷ The band between 2800 and 3000 cm^{-1} shows the presence of C–H stretching vibrations. Absorption bands between 675 – 900 cm^{-1} ¹⁶⁸ and 1575 – 1625 cm^{-1} indicate the

presence of monocyclic, polycyclic and substituted aromatic groups.⁶⁹ C=C stretching vibrations, which indicate the presence of alkenes, are confirmed by the absorbance band between 1625 and 1675 cm^{-1} .⁶⁵ The band between 850 – 950 cm^{-1} for SD shows the presence of ethers. In comparison to the biochar, the band between 950 and 1325 cm^{-1} most probably represents the presence of primary, secondary and tertiary alcohols and phenols as a result of the C–O stretching and O–H in plane distortion of these functional groups.⁶⁶ The alcohol/phenol OH functional group was no longer visible at 3300 cm^{-1} , indicating that phenolic biopolymer, which is characteristic of lignin in the digestate, has been removed. This means that the bio-aromatic composition with the potential to be valorised to biofuels has been burned off.

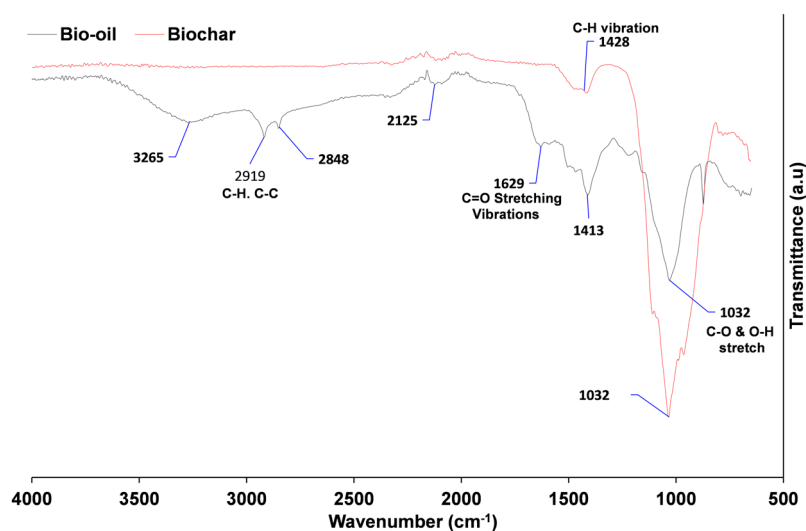


Fig. 3 Fourier transform infrared (FTIR) spectrum of bio-oil and Biochar in the range of 500 – 4000 cm^{-1} .



However, the C–C was modified to an inflection at $\sim 1600\text{ cm}^{-1}$, which is characteristic of an alkene C=C.⁶⁴

Activity of catalysts effect on bio-oil yield, conversion and selectivity from SD

Product yield (aromatic and aliphatic) from hydrogenolysis depolymerisation was based on substrate type. Product yield based on substrate conversion seems to be higher with nickel on zirconium and rhodium on carbon than copper on alumina, palladium and ruthenium on a carbon support. Fig. 4 reports the bio-oil yields for Ni/ZrO₂ (13.43 wt%), Rh/C (12.40 wt%), Cu/Al₂O₃ (10.92 wt%), Pd/C (7.45 wt%) and Ru/C (6.92 wt%), which indicates that the order of performance (based on bio-oil yield) was Ni/ZrO₂ > Rh/C > Cu/Al₂O₃ > Pd/C > Ru/C. Experiments for bio-oil yield were repeated twice; the results were reproducible with a small standard deviation (Table S1, ESI†), and so data is reported with the experimental error bars of $\pm 4.97\%$ (calculation shown in ESI†).

Fig. 5 reports the substrate conversion based on the three most active catalysts (Ni/ZrO₂, Rh/C and Cu/Al₂O₃) as a function of time. When the reaction time was maintained for 1 h, the substrate conversion over Ni/ZrO₂, Rh/C, and Cu/Al₂O₃ was 40.30, 38.08 and 36.04%, respectively. For 2 h reaction time, conversion over Ni/ZrO₂, Rh/C, and Cu/Al₂O₃ was (37.40, 34.11, and 33.12%, respectively); 3 h reaction time, conversion was (33.05, 31.50 and 30.80%, respectively); 4 h reaction time, conversion was (32.01, 31.20 and 30.10%, respectively). The substrate conversion trend for all the catalysts showed that catalytic activity seems to be highest in the 1 h reaction; in the order, 1 h > 2 h > 3 h > 4 h. Given the decrease in the yields of bio-oil with time as reported, it might be expected that the biochar yield should have increased and be highest after 4 h.

In the absence of head space analysis a possible explanation is that this is due to further production of volatiles as the reaction progressed. It is proposed that the drop in yield at longer run times is due to some of the liquid products are being converted slowly by the catalyst to volatile, gaseous products that are not being isolated at the end of the reaction. Such losses could be minimised by stopping the reaction sooner.

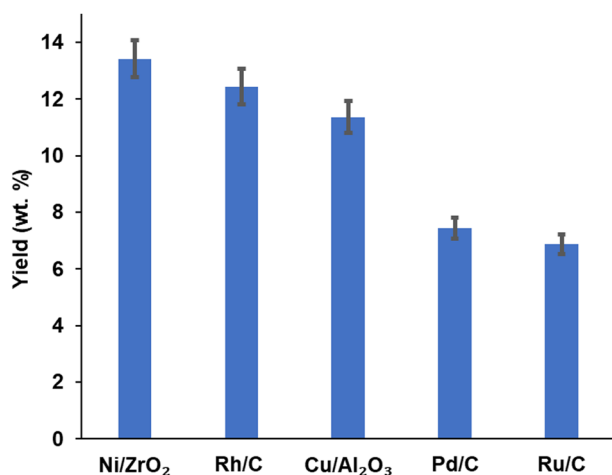


Fig. 4 Affect of catalyst selection on bio-oil yield.

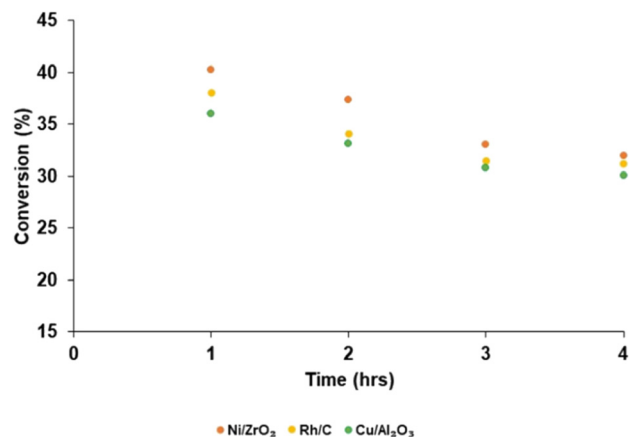


Fig. 5 Substrate conversion as a function of reaction time for different catalysts.

Even if there are more moles of biochar produced it may well weigh less than the substrate due to dehydration/elimination of volatile components. As such, conversion of the substrate may well have continued in this time. However, since the bio-oil yield also decreased over this period, although further conversion of the substrate to biochar cannot be discounted entirely, the only way to achieve a mass balance is to consider that more and more volatile products are produced (see Fig. S2–S6 in ESI† for biochar yield).

Fig. 6 shows the catalyst selectivity (*S*) toward bio-oil at different reaction times (1 h, 2 h, 3 h and 4 h). Selectivity increases as the reaction time increases with the Ni/ZrO₂ catalysts; for a reaction run time of 1 h, product selectivity was 33.0%. When the reaction was run for 4 h, selectivity increased to 35.6%. This trend was different for Rh/C when the reaction run time was 1 h; selectivity was 32.64% but reduces to 30.3% for a 4 h run time reaction. Also, for Cu/Al₂O₃, selectivity was 30.3% when the reaction was run for 1 h; it decreases when the reaction time was for 2 h.

However, when the reaction was run for 3 h; product selectivity increased to 31.0%. The bio-oil yield depends not only on catalytic activity and selectivity but also on the biomass substrates and the extraction solvent. It is believed that the

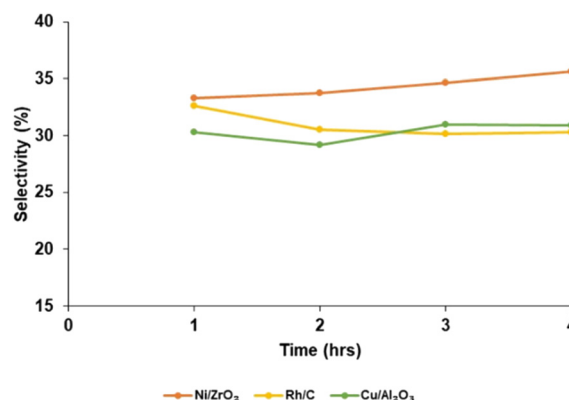


Fig. 6 Product selectivity as a function of reaction time for different catalysts.



effectiveness of nickel–metal in cleaving both the C–O and β -O-4 bonds,^{39,70} as well as the amphoteric^{49,68} characteristic of ZrO₂ support, plays a major role. The redox property of nickel is influenced by the acid/base properties of the support.^{51,71} Also, the affinity of rhodium to the β -O-4, which is the predominant ($\sim$ 50%) linkage in lignin, is higher than the affinity of the precious metals on the other studied heterogeneous catalysts. Extracting solvent also plays a major role in bio-oil yield, for instance, in a previous study⁵⁰ with Rh/C and Pd/C using cork biomass substrate and dioxane as extracting solvent, the bio-oil yield was 11.50 and 7.2 wt%, respectively. Using 2-MeTHF solvent in this current work indicates a 7.82 and 3.47% increase for Rh/C and Pd/C, respectively. Furthermore, higher bio-oil yield can be achieved by the addition of base (BCD) to the solvent as investigated and reported by McCallum *et al.*⁵¹

Ma *et al.*³⁹ reported a 15.1 wt% yield from organosolv lignin depolymerisation through hydrogenolysis using Ni/ZrP. Schutyser *et al.*⁵² reported yields of above 80% using models (lignin-derived guaiacols). Van den Bosch *et al.*⁷² reported the delignification of birch sawdust using a Ru/C catalyst *via* the simultaneous solvolysis and hydrogenolysis with a lignin-oil yield of above 50% phenolic monomers and about 20% phenolic dimers. It was said that the higher selectivity to phenolic monomers with Ru/C was as a result of its affinity for the ether-bonds between phenolic units that prevents re-polymerisation reactions, hence, a stable C–C bonds within the lignin structural network.

GC–MS and NMR characterisation of bio-oil

Proposed mechanism and reaction pathway. The production of smaller molecules from lignin is actualised *via* depolymerisation,

which is obtained by the cleavage of the linkages between the phenyl propane units.^{32,70} Typical bonds present in lignin are: diphenyl ether (β -O-4, which constitute above 50% of the bonding structures as shown in Fig. 7), 5–5, β -5, α -O-4, β - β , β -1, 4-O-5 and dibenzodioxocin.⁷³ The major pathway in which lignin is depolymerised is through the cleavage of the ether bond. Cleavage of this linkage leads to the formation of water-soluble compounds consisting of phenolic hydroxyl groups due to fragmentation. The disruption of the β -O-4 linkage forms simpler model compounds that look like *p*-coumaryl, coniferyl and sinapyl monolignols.⁷⁴ Reductive delignification uses a redox catalyst in combination with H₂ or an H-donor to cleave the inter-unit ether bonds (β -O-4, α -O-4) and side-chain hydroxyl groups in lignin. The proposed mechanisms and pathways for lignin reductive depolymerisation result in three routes: (i) hydrogenolysis of the ether bonds, (ii) removal of benzylic OH-groups (OH α), and (iii) removal of OH γ -groups.^{75,76}

Using the GC–MS to analyse the composition of the derivatised bio-oil produced from the SD hydrogenolysis catalysed reaction, the major product extracted was ethylguaiacol (which is a bio-aromatic oxygenate octane booster). This product was obtained from the cleavage of aryl glycerol ether (β -O-4) and diphenyl ether (5-O-41) linkages in the lignin structure, as shown in ESI† (Fig. S1). The mass spectrum of TMS-derivatised ethylguaiacol is shown in ESI† (Fig. S7). In previous studies, Yueyuan⁷⁷ reported the extraction of ethylguaiacol from the hydrogenolysis of lignosulphonate. Garrett *et al.*⁵⁰ and Graca *et al.*⁷⁸ also reported aromatic species extracted from barks as propylguaiacol, dihydroconiferylalcohol and other aromatics

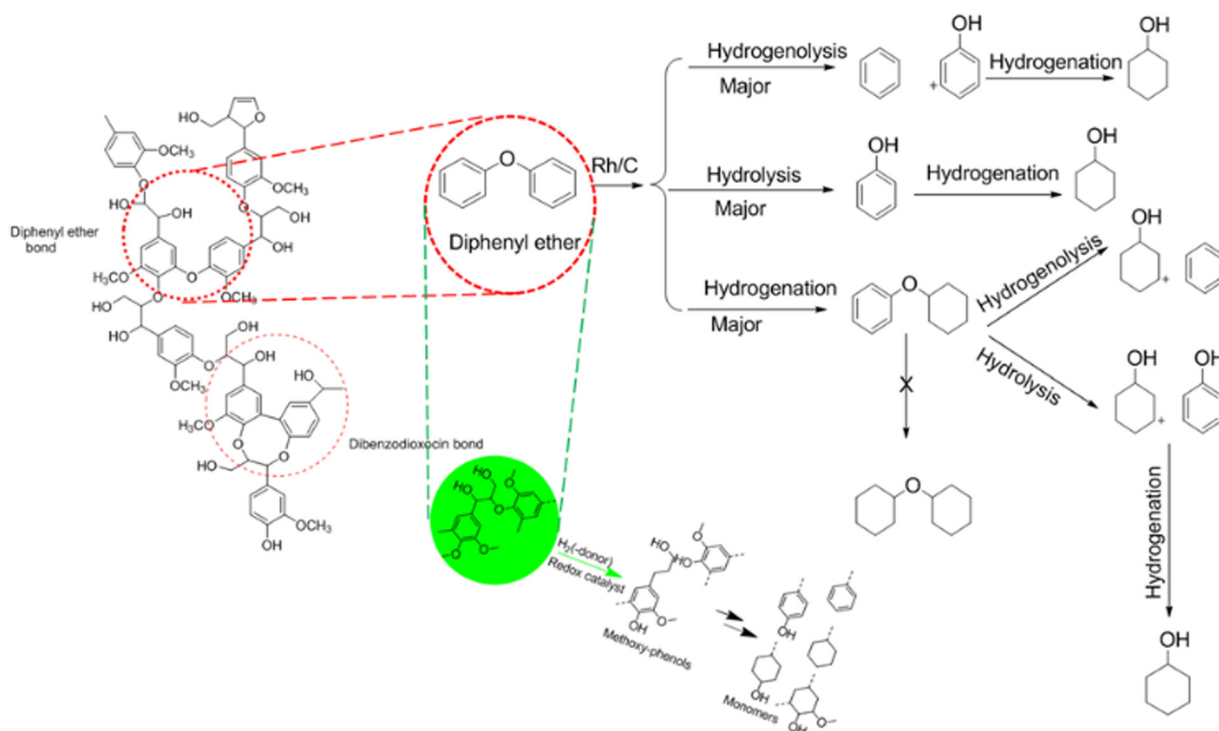


Fig. 7 The proposed mechanism and pathway lignin reductive depolymerisation routes.

such as catechol, ferulic acid and 3,4-dihydroxybenzoic acid in small amounts.

^1H NMR spectroscopy was performed and used to determine the quantitative analysis of the isolated crude bio-oil. The side-chain methylene signals α to the aromatic rings of ethylguaiacol. This analysis showed the presence of both fatty acid and aromatic products. The multiple peaks could potentially be as a result of the partial depolymerisation of the lipids and esters. The side-chain methylene signals α to the aromatic rings of ethylguaiacol. In the region between 6.4 and 7.0 ppm, the aromatic hydrogens on the guaiacyl substituent are evident. Quantification of the total lipid and fatty acid content was determined using a vanillin internal standard and comparing the integration of aldehydic singlet in the ^1H NMR spectrum at 9.73 ppm to the distinctive peak at 2.2–2.5 ppm from the CH_2 group alpha to the ring as the fatty acid/ester carbonyl group. The NMR peak integration was used to calculate the percentage of the aromatic products in the bio-oil using the formula in the ESI† (determination of aromatic yields). The ^1H NMR spectrum is reported in Fig. 8 and ESI† (S8); Table 2 shows the summary.

Characterising the most active catalyst

To understand the surface morphology of the most active catalyst herein SEM-EDX mapping along with TEM analyses were performed. TEM images in Fig. 9(a) and (b) showed spherical particles with a particle size of ~ 9 nm. The nickel-based catalyst, which is denoted as Ni/ZrO₂, showed uniform particle size distribution, as shown in Fig. 9(c).

The elemental mappings of Ni, Ce, Zr, Si, Al and O elements show the uniform distribution of Ni active sites of the surface of the Ni-based catalyst as shown in Fig. 9(d)–(j) while Al mapping shows a small cluster formation as shown from the circle in Fig. 9(d) and (i). The presence of oxygen carrier in the catalyst

Table 2 Calculation of aromatic yields

Substrate/catalyst	Run 1		Run 2		Ref.
	Yield (mg)	Aromatics (wt%)	Yield (mg)	Aromatics (wt%)	
Rh/C					
Cork ^a	40.0	2.0	46.0	2.1	50
Spruce ^a	36.0	1.2	45.0	1.5	50
Sycamore ^a	114.0	3.8	81.0	2.7	50
SD ^b	250.8	8.4	210.5	7.0	This work
Pd/C					
Cork ^a	26.0	1.3	38.0	1.9	50
Spruce ^a	45.0	1.5	63.0	2.1	50
Sycamore ^a	63.0	2.1	46.0	1.9	50
SD ^b	150.6	5.0	120.3	4.9	This work

^a Hydrogenolysis with dioxane solvent and reaction time at 4 h.

^b Hydrogenolysis with MeTHF solvent and reaction time at 1 h. Distilled water : 2-MeTHF is 1 : 1.

composition in the form of CeO₂, ZrO₂ along with SiO₂ and Al₂O₃ will aid in the oxidation of any carbon deposition over the surface of the Ni-based catalyst during the reaction.^{79,80}

Comparison with other work

The weight percentages of the aromatic in the bio-oil reported in this work for the first and second experimental run using Rh/C catalysts were 8.4 and 7.0 wt% respectively. When Pd/C was used, the percentage aromatic for the first and second experimental run was 5.0 and 4.9 wt% respectively. These values compare favourably with other work as compared in Table 2, although in this work, the selectivity to aromatic is higher as compared to previous work using dioxane solvent.⁵⁰ Also, McCallum *et al.*⁵¹ reported the weight percentage of the aromatic in bio-oil extracted from cork using different ratios of distilled water to 2Me-THF. Using distilled water to 2-MeTHF ratios of 1 : 9, 4 : 6, 6 : 4 and 9 : 1, the reported weight percentage of aromatic in the bio-oil was 3.5, 7.1, 4.7 and 5.1 wt%, respectively.

Life cycle assessment

Inventory analysis: Previous studies (*e.g.* Girio *et al.*,⁸¹) have focused on the relative CO₂ equivalent production between different processes and the transport of biomass to the processing site. In this study, the focus is on the net energy balance between the inputs of the process and the outputs in the form of the bio-oil distillate. The transportation distance between the anaerobic digestion plant and the treatment unit was conservatively assumed as 20 km. The energy requirements for transportation were assumed to be 5.7 MJ tkm⁻¹.^{82,83}

After the transportation of anaerobic digestate, impacts were considered for the preparation of feedstock for the hydrogenolysis reaction. Although lab-scale preparation used oven drying in this experiment, it was assumed that for large-scale commercial operations, a decanting centrifuge and belt dryer would be employed for producing solid digestate.

A decanting centrifuge can dewater anaerobic digestate to produce a solid fraction using electrical energy as input

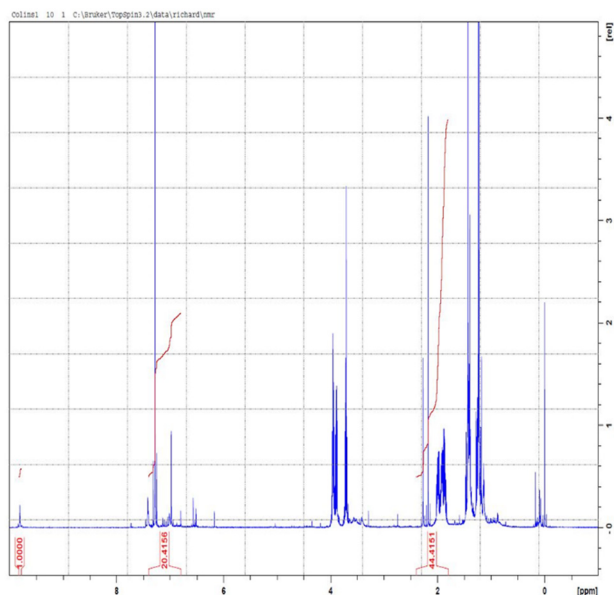


Fig. 8 ^1H NMR spectrum of Me-THF extracted product from SD hydrogenolysis.



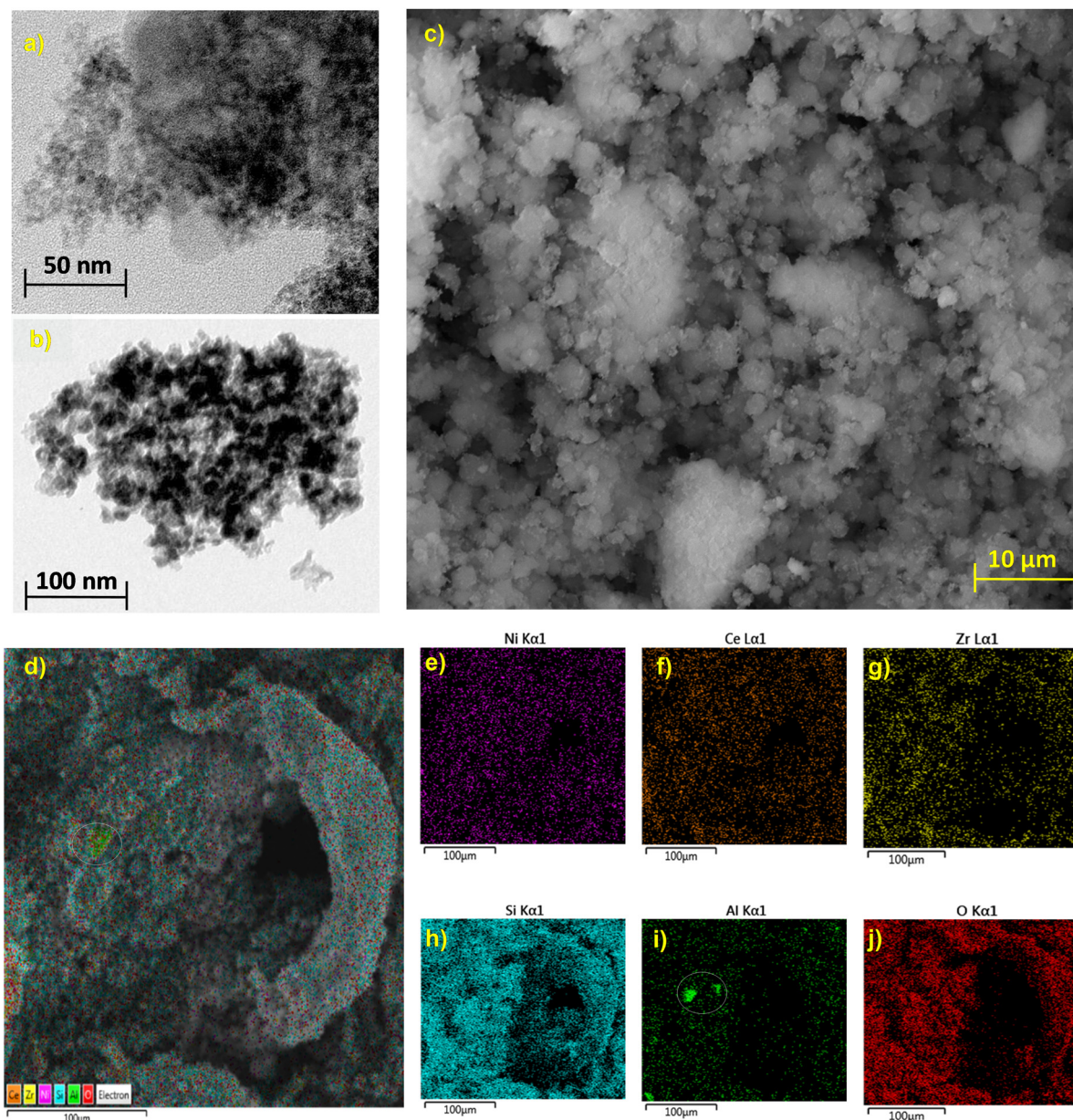


Fig. 9 Surface morphology characterisations of Ni/ZrO₂; TEM images (a), (b), SEM image (c) and elemental mapping (d)–(j).

(3.5 kW h per tonne of anaerobic digestate dewatered).⁸⁴ It was assumed that the total solids of the anaerobic digestate was 6.3%. The separation efficiency of a decanting centrifuge is shown in Table S2 of the ESI.† For one tonne of anaerobic digestate treated, 126 kg of solid fraction production was assumed. This solid fraction, when treated *via* belt dryer, can produce 36 kg of solid digestate (at 10% moisture) with thermal energy as input for raising the temperature and evaporating moisture (total energy input of 2.63 MJ kg^{−1} of solid fraction).

Finally, for the hydrogenolysis reaction procedure (mentioned in the Methods Section), it was calculated that 6 kg of Ni/ZrO₂ catalyst, 225 kg of 2 Me-THF and 264 kg of deionised water would be required for the conversion of 36 kg of solid digestate. Calculations were performed for the Ni/ZrO₂ catalytic

system as it showed the highest bio-oil yield. The electrical energy required for hydrogenolysis was taken from Yano *et al.*⁸⁵ as 0.16 kW h kg^{−1} of feedstock treated (Table 3). The amount of bio-oil produced at 17% yield was 90.34 kg, which is equivalent to 1445 MJ of energy based on a calorific value of 16 MJ kg^{−1}.⁸⁶

Energy balance: The total energy required for the process was calculated as 849 MJ. The energy consumed for the hydrogenolysis reaction stage was highest at 391 MJ (46%), closely followed by the digestate dewatering and drying stage (344 MJ) (41%). The total output energy in the form of bio-oil was 1445 MJ. The net energy is defined as the ratio of output energy to input energy and was thus calculated as 1.7 (a net energy ratio > 1 shows that there was net production of energy during the process). This is over double the net energy of petroleum-



Table 3 Inventory data for conversion of 1 tonne of anaerobic digestate to bio-oil

Input/process	Unit	Input	Output
Transportation			
Feedstock transportation	tkm		20
Anaerobic digestate	t	1	
Energy (fossil fuels) ^a	MJ	114	
Solid-liquid separation (decanter centrifuge)			
Digestate	t	1	
Electrical energy	MJ	12.6	
Liquid fraction (disposed)	kg		874
Solid fraction	kg		126
Drying of solid fraction (belt dryer)			
Solid fraction	kg	126	
Thermal energy	MJ	331.2	
Produced solids (10% moisture)	kg		36
Hydrogenolysis			
Produced solids (10% moisture)	kg	36	
Ni/ZrO ₂ catalyst ^b	kg	6	
2 Me-THF ^c	kg	225.4	
Deionised water ^d	kg	264	
Electrical energy	MJ	20.8	
Bio-oil	kg		90.3

^a Freight transport by lorry, 3.5–7.5 metric tonnes, euro 6.⁸² ^b Due to a lack of data on production of Ni/ZrO₂ catalysts, it was considered that a reaction between Ni and ZrO₂ would lead to production of Ni/ZrO₂ (100% yield assumed). For production of 1 kg of Ni and ZrO₂, energy requirements were adapted from Nuss *et al.*⁸³ ^c Due to a lack of data on the production of 2Me-THF, it was considered that a reaction of furfural (1.11 kg) and hydrogen gas (82.3 g) would be needed for 1 kg of 2Me-THF (100% yield assumed). The energy requirement for furfural production was sourced from Raman and Gnansounou.⁸⁷ ^d De-ionised water, reverse osmosis, production mix, at plant, from groundwater.⁸²

based petrol and diesel⁸⁸ and is comparable to conventional temperate crop-based biofuels.⁸⁹

Conclusion

In this study, we report the valorisation of SD from AD *via* hydrogenolysis reaction using both heterogeneous platinum group and active metal catalysts to produce phenolic aromatic monomers from the lignin constituent of the SD. The major product extracted was ethylguaiaicol. The bio-oil yield ranged between 6.92–13.43 wt%, with Ni/ZrO₂ giving the highest yield (13.43 wt%) and Ru/C giving the lowest yield (6.92 wt%). The effect of using a green environmental benign solvent (2-methyltetrahydrofuran) was evident as the yield increased by ~17 wt% as compared to the use of dioxane (11.5 wt%) as a reaction solvent with Ni/ZrO₂ catalysts. The life cycle assessment was performed considering one tonne of anaerobic digestate converted to biofuels as feedstock. System boundary included transportation of digestate, dewatering and drying of digestate, hydrogenolysis reaction of solid digestate in presence of Ni/ZrO₂ catalyst. The results showed that a total of 849 MJ of input energy would be required for the conversion for one tonne of digestate to bio-oil, resulting in an energy ratio that is similar to conventional biofuels and over double that of fossil transport fuels.

Disclaimer

The views and opinions expressed in this paper do not necessarily reflect those of the European Commission or the Special EU Programmes Body (SEUPB).

Conflicts of interest

There are no conflicts to declare.

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