UNIVERSITY^{OF} BIRMINGHAM University of Birmingham Research at Birmingham

A Kinetic Model of Furfural Hydrogenation to 2-Methylfuran on Nanoparticles of Nickel Supported on Sulfuric Acid-Modified Biochar Catalyst

Mudi, Ismaila; Hart, Abarasi; Ingram, Andy; Wood, Joe

DOI: 10.3390/catal14010054

License: Creative Commons: Attribution (CC BY)

Document Version Publisher's PDF, also known as Version of record

Citation for published version (Harvard):

Mudi, I, Hart, A, Ingram, A & Wood, J 2024, 'A Kinetic Model of Furfural Hydrogenation to 2-Methylfuran on Nanoparticles of Nickel Supported on Sulfuric Acid-Modified Biochar Catalyst', *Catalysts*, vol. 14, no. 1, 54. https://doi.org/10.3390/catal14010054

Link to publication on Research at Birmingham portal

General rights

Unless a licence is specified above, all rights (including copyright and moral rights) in this document are retained by the authors and/or the copyright holders. The express permission of the copyright holder must be obtained for any use of this material other than for purposes permitted by law.

•Users may freely distribute the URL that is used to identify this publication.

•Users may download and/or print one copy of the publication from the University of Birmingham research portal for the purpose of private study or non-commercial research.

•User may use extracts from the document in line with the concept of 'fair dealing' under the Copyright, Designs and Patents Act 1988 (?) •Users may not further distribute the material nor use it for the purposes of commercial gain.

Where a licence is displayed above, please note the terms and conditions of the licence govern your use of this document.

When citing, please reference the published version.

Take down policy

While the University of Birmingham exercises care and attention in making items available there are rare occasions when an item has been uploaded in error or has been deemed to be commercially or otherwise sensitive.

If you believe that this is the case for this document, please contact UBIRA@lists.bham.ac.uk providing details and we will remove access to the work immediately and investigate.





Article A Kinetic Model of Furfural Hydrogenation to 2-Methylfuran on Nanoparticles of Nickel Supported on Sulfuric Acid-Modified Biochar Catalyst

Ismaila Mudi¹, Abarasi Hart², Andrew Ingram¹ and Joseph Wood^{1,*}

- ¹ School of Chemical Engineering, University of Birmingham, Edgbaston, Birmingham B15 2TT, UK; ixm928@student.bham.ac.uk (I.M.); a.ingram@bham.ac.uk (A.I.)
- ² Energy and Bioproducts Research Institute, Aston University, Birmingham B4 7ET, UK; a.hart1@aston.ac.uk
- * Correspondence: j.wood@bham.ac.uk

Abstract: Lignocellulosic biomass can uptake CO_2 during growth, which can then be pyrolysed into three major products, biochar (BC), syngas, and bio-oil. Due to the presence of oxygenated organic compounds, the produced bio-oil is not suitable for direct use as a fuel and requires upgrading via hydrodeoxygenation (HDO) and hydrogenation. This is typically carried out over a supported metal catalyst. Regarding circular economy and sustainability, the BC from the pyrolysis step can potentially be activated and used as a novel catalyst support, as reported here. A 15 wt% Ni/BC catalyst was developed by chemically modifying BC with sulfuric acid to improve mesoporous structure and surface area. When compared to the pristine Ni/BC catalyst, sulfuric activated Ni/BC catalyst has excellent mesopores and a high surface area, which increases the dispersion of Ni nanoparticles and hence improves the adsorptive effect and thus catalytic performance. A liquid phase hydrogenation of furfural to 2-methylfuran was performed over the developed 15 wt% Ni/BC catalyst. Langmuir-Hinshelwood-Hougen-Watson (LHHW) kinetic type models for adsorption of dissociative H₂ were screened based on an R^2 value greater than 99%, demonstrating that the experimental data satisfactorily fit to three plausible models: competitive (Model I), competitive at only one type of adsorption site (Model II), and non-competitive with two types of adsorption sites (Model III). With a correlation coefficient greater than 99% between the experimental rates and the predicted rate, Model III, which is a dual-site adsorption mechanism involving furfural adsorption and hydrogen dissociative adsorption and surface reaction, is the best fit. The Ni/BC catalyst demonstrated comparative performance and significant cost savings over previous catalysts; a value of 24.39 kJ mol⁻¹ was estimated for activation energy, -11.43 kJ mol⁻¹ for the enthalpy of adsorption for H_2 , and $-5.86 \text{ kJ mol}^{-1}$ for furfural. The developed Ni/BC catalyst demonstrated excellent stability in terms of conversion of furfural (96%) and yield of 2-methylfuran (54%) at the fourth successive experiments. Based on furfural conversion and yield of products, it appears that pores are constructed slowly during sulfuric acid activation of the biochar.

Keywords: furfural; sulfuric acid activation of biochar; hydrogenation; kinetic modelling

1. Introduction

As part of climate change mitigation, fossil fuel transportation demands can be substituted with biomass-derived biofuels. The development and use of biomass resources, as well as platform chemicals derived from biomass, has been considered an alternative substitute to fossil fuels in the face of increasing energy demands and climate change [1]. Due to its carbon neutrality viewpoint, ecological, and economic sustainability, renewable biomass has attracted considerable attention for conversion into biofuels and chemicals. In the pursuit of sustainable alternatives to petroleum-based fuels and chemicals, it is becoming more important to convert lignocellulosic biomass into biofuels and platform chemicals. Furfural, a platform chemical obtained from biomass, may be further processed to provide



Citation: Mudi, I.; Hart, A.; Ingram, A.; Wood, J. A Kinetic Model of Furfural Hydrogenation to 2-Methylfuran on Nanoparticles of Nickel Supported on Sulfuric Acid-Modified Biochar Catalyst. *Catalysts* **2024**, *14*, 54. https:// doi.org/10.3390/catal14010054

Academic Editor: Xiaofeng Wang

Received: 12 December 2023 Revised: 8 January 2024 Accepted: 9 January 2024 Published: 11 January 2024



Copyright: © 2024 by the authors. Licensee MDPI, Basel, Switzerland. This article is an open access article distributed under the terms and conditions of the Creative Commons Attribution (CC BY) license (https:// creativecommons.org/licenses/by/ 4.0/). furfuryl alcohol, 2-methylfuran, furan, tetrahydrofurfuryl alcohol, cyclopentanone, and tetrahydrofurfuryl alcohol [2–5]. As one of the most promising lignocellulosic biomass platform molecules, furfural is derived from hemicellulose by hydrolysing and dehydrating xylose, which is C₅ sugar [6,7]. The global production of furfural is almost 5,000,000 tons per annum from a variety feedstocks [8], and it is also a typical oxygenated model compound found in biomass-derived bio-oils. Consequently, biomass-derived compounds contain a high level of oxygen, so selective hydrodeoxygenation (HDO) and hydrogenation have become a primary objective in most research studies.

It is difficult to control the selectivity of the products in furfural hydrogenation because of many possible reactions; for example, hydrogenolysis of C=O bonds, decarbonylation, and hydrogenation of C=O bonds or furan rings [9,10]. Catalytic hydrogenation of furfural to 2-methylfuran can be performed in either the gas or the liquid phase. In comparison with gas-phase hydrogenation of furfural, which requires higher energy consumption due to vaporisation and the harsh reaction temperature that often results in catalyst deactivation [11], liquid-phase hydrogenation needs only a suitable temperature for reaction. As such, it would be advantageous to develop a solid catalyst capable of hydrogenating liquid furfural efficiently and selectively into 2-methylfuran.

Several value-added products can be produced from furfural that has the furan ring and the –CHO group which are unsaturated C=O and C=C bonds. In the production of 2-methylfuran, the furanic "O" and the double bond of the ring are not affected by selective hydrodeoxygenation (HDO). This involves the selective removal of the oxygen from the C=O side group of the furan while suppressing the competitive ring hydrogenation [6,10]. Through hydrogenolysis of C=O bonds, furfural undergoes HDO into 2-methylfuran. Thus, catalyst design and process optimisation can be challenging when it comes to selectively producing 2-methylfuran during a multiproduct reaction of furfural hydrogenation. Hence, a catalyst candidate for this conversion would possess catalytic active sites that are capable of excellent hydrogenation and hydrogenolysis. It is essential to develop a catalyst with appropriate activity, selectivity, and stability to achieve significant selectivity toward 2methylfuran. The chemical compound 2-methylfuran is crucial to the production of furan compounds, pentanediol, important intermediate for pharmaceuticals, fuel additive, etc. It can also be partly blended with gasoline for engines and has a high octane number, boiling point, and energy density [6,12].

Catalytic hydrogenation of furfural to produce either 2-methylfuran or furfuryl alcohol in liquid phase had been explored using the following catalysts: Pt/g-C₃N₄, graphitic carbon nitride nanosheet [12], NiCoZn@CN [13], Pt/SiO₂, Pt/ZnO, Pt/MgO, Pt/y-Al₂O₃, and Pt/CeO₂ [14], Pt–Sn/SiO₂ [4], CuZnAl hydrotalcite-like (HTLc) catalyst [11], Ir/C [8], Cu/ZrO₂, Pd/ZrO₂, and Cu-Pd/ZrO₂ [12], Mo₂C [6], Cu/SiO₂, Cu/Al_2O_3 , and Cu/ZnO[1], and $Cu-Co/SiO_2$, Cu-Co/H-ZSM5, and $Cu-Co/\gamma-Al_2O_3[10]$. Consequently, Pd/C, Pt/C, Re/C, Ru/C, Rh/C, Ni/C, and Cu/C catalysts have been used to study the hydrogenation, HDO, and ring opening of furfural [15]. Also, comprehensive reviews have been published by several authors that discusses recent advances in the catalytic hydrogenation of furfural towards furfuryl alcohol, 2-methylfuran, furan, and other value-added products in terms of different noble metals and non-noble metals catalysts, reaction mechanisms related to the different catalytic materials, and reaction conditions [5,16]. However, most studies focus on catalysts that are selective toward furfuryl alcohol product. On one hand, the chemisorption of aromatic compounds and produced water from HDO reactions usually deactivate most of these support materials [17]. Noble metals and bimetallic catalysts, on the other hand, are prohibitively expensive coupled with deactivation, limiting their industrial scale and widespread practical applicability. Thus, hydrogenation and HDO of oxygenate model compounds of bio-oil could become more cost effective and sustainable if biochar (BC), which is one of the by-products of biomass pyrolysis, can be recycled for catalyst support. It is also important to note that pristine BC, which is derived from biomass via pyrolysis, has certain constraints in the pore structure [18], which limits mass transport during reactions. Thus, catalysis may be

improved by synthesising highly mesoporous BC as catalyst support by increasing surface area and pore structure, which improves active metal dispersion. Studies on BC revealed that it has a relatively moderate porosity and surface area and possesses mineral elements such as K, Ca, and P, and functional groups on the surface [19,20].

In general, hydrogenation, HDO, and hydrogenolysis of the phenol, aldehyde, and methoxy groups of typical bio-oil model compounds are possible with the application for Ni-based catalysts. Therefore, a catalyst for furfural hydrogenation is developed using nickel and BC combined. Wet impregnation was used to immobilise Ni nanoparticles on mesoporous BC support. In this study, the mesoporosity of BC is enhanced through chemical modification using an acidic solvent, sulfuric acid (H_2SO_4). Through sulfuric acid treatment, the BC support material is tuned to improve the catalyst's textural attributes, including surface area, mesoporosity, and pore volume. The objective is to enhance mass transport of reactants and increase specific surface area to improve the dispersion of nickel nanoparticles, and consequently catalyst performance. In our previous work, using Fouriertransform infrared spectroscopy (FTIR) technique, it was found that the surface of the BC contains functional groups such as the hydroxyl OH-group, carboxylic acids -COOH, -C-O stretching of cellulose, and C=C of aromatic systems [21]. These surface functional groups could help improve metal-support interaction. To model the kinetics of this catalyst, a Langmuir–Hinshelwood–Hougen–Watson (LHHW) kinetic model for hydrogenation or HDO of bio-oil model compounds has been proposed for different catalysts based on earlier research works; e.g., vanillin, 1-heptyne and furfural substrates were applied [22–25]. These assume a single or dual nature of adsorption sites and a surface reaction between a partly hydrogenated furfural species and a dissociative adsorbed hydrogen. Using a batch reactor, 2-methylfuran is produced by liquid-phase catalytic hydrogenation of furfural utilising tetralin as a solvent over nickel supported on sulfuric acid modified biochar catalysts. The objective of this work is to develop a kinetic model of furfural hydrogenation into 2-methylfuran bespoke to the prepared Ni/BC catalyst. Thus, bio-oil catalytic upgrading with Ni/BC includes significant cost savings over previous catalysts, promotes circular economy, and is sustainable with a comparative performance to previous catalysts. The novelty of the study lies in the development of renewable and cost-effective catalysts for hydrodeoxygenation and hydrogenation of bio-oils. The construction of more mesopores within biochar by acid activation is needed to improve surface area, enhance metal nanoparticle dispersion, and increase catalytic activity and selectivity. Thus, circular economy and sustainability in the field of bio-oil upgrading can be achieved by activating biochar from the pyrolysis step and using it as a catalyst support. This study shows that the developed Ni/BC catalyst provides a cost effective and efficient means of hydrogenating furfural into 2-methylfurfuran.

2. Results and Discussion

A comparison of porosimetry properties of biochar after sulfuric acid treatment is shown in Table 1. As a result of sulfuric acid treatment, the biochar gains more microporous (less than or equal to 2 nm) and mesoporous (2 nm to 50 nm) pores, and an increase is observed in pore volume for pore sizes greater than 5 nm, 2–5 nm, and smaller than 2 nm. In sulfuric acid-treated biochar, there are 7 times more micropores than in the biochar received, while the number of mesopores is 3.2 times greater. This suggests that sulfuric acid creates more micropores and mesopores as a result of the reaction with O-containing species in the biochar. Due to this, sulfuric acid-treated biochar has 4.7 times the surface area of as-received biochar. However, the mean pore diameter of as-received biochar is higher than that of sulfuric acid-treated one. The modified biochar is identified as Type IV isotherm based on the adsorption–desorption hysteresis patterns.

In previous study, a detailed characterisation of Ni/BC catalysts was reported [21]. The developed Ni/BC catalyst characteristics reported already include pore size distribution based on nitrogen adsorption technique using Barrett–Joyner–Halenda (BJH) equation; transmission electron microscopy (TEM) report on the distributions of Ni nanoparticles (between 4 nm and 16 nm) over biochar support (Figure S1); X-ray diffraction (XRD) patterns of the biochar and Ni/BC that revealed characteristic peaks of NiO at 20 values of 37° , 44.5°, and 64.5°, respectively. Furthermore, a Fourier-transform infrared (FTIR) spectrometer was used to identify a number of complementary functional groups, including a phenolic OH-group at 3400 cm⁻¹ and carboxylic acids, –COOH, at 2330 cm⁻¹ (Figure S2). Additionally, a reduction peak corresponding to the reduction of the impregnated Ni metal and its oxides (NiO) in the catalyst was identified at temperatures between 300 °C and 450 °C using hydrogen temperature-programmed reduction (H₂-TPR); the number and strength of the active sites of the Ni/BC catalyst were determined using carbon dioxide

temperature-programmed desorption (CO₂-TPD) (Figure S3); and thermogravimetric analysis was used to examine the catalysts' thermal stability. These characterisation results were published in the literature [21]. The amount of Ni metal incorporated into the biochar was determined using inductively coupled plasma-optical emission spectroscopy (ICP-OES). According to ICP-OES analysis results, the developed Ni/BC catalyst had a metal loading of 14.7 wt% (Table S1). Details can be found in Supplementary Information.

Table 1. Changes in biochar porosimetry properties due to sulfuric acid treatment.

Parameter	As-Received Biochar	H ₂ SO ₄ -Modified Biochar		
Pore Size (nm)	Pore Volume (cm ³ g ⁻¹)			
≥5	0.0595	0.1199		
2–5	0.0175	0.1281		
<u>≤2</u>	0.0043	0.0302		
	Specific surface area (m ² g ^{-1})			
_	74.84 ± 1.05	353.50 ± 3.68		
Total pore volume (cm 3 g $^{-1}$)	0.0813	0.2781		
Average pore size (nm)	5.791	4.322		

2.1. Mass Transport Study

Performance of slurry reactors may be affected by reaction kinetics as well as resistance to gas absorption and mass transport at catalyst particles. Prior to surface catalytic reaction in a slurry reactor (i.e., gas-liquid-solid phase system), several mass transport processes must be taken into consideration, such as gas-liquid mass transfer, liquid-solid mass transfer, and internal pore diffusion combined with surface chemical reactions [26]. The elimination of transport restrictions, which include impacts of external and internal heat and mass transfer as well as the rate of hydrogen transfer from the gas phase to the liquid phase, is a crucial step in the process of validating quantitative kinetic models [27]. To guarantee that the data are gathered under kinetically controlled reaction regime, the furfural hydrogenation reaction is carried out in this study under conditions where both external and internal mass transport restrictions are insignificant, as necessary for kinetic model development. Furthermore, due to the tendency of the catalyst to settle, it may be difficult to mix and disperse the catalyst within the solvent depending on catalyst concentration, average particle size, and stirring rate [28,29]. Adequate agitation is necessary for maintaining a uniform concentration and temperature throughout the reactor medium, but it must also be sufficient for mass transfer across the thickness of the boundary layer surrounding the catalyst particles [22,30]. To ensure the experiments are performed under negligible external mass transfer restrictions, the conversion and 2-methylfuran yield from furfural hydrogenation over the Ni/BC catalyst were examined as a function of mixing, with stirring speeds ranging from 600 to 1200 rpm. Figure 1 shows the conversion of furfural and the yield of 2-methylfuran as a function of the stirring rate. Notably, the furfural conversion increases as stirring rate increases from 600 rpm to 800 rpm, and then plateaus between 800 rpm and 1200 rpm (Figure 1a), whereas the yield of 2-methylfuran,



and the corresponding rate of furfural hydrogenation, increases as the mixing speed rate increases from 600 rpm 1000 rpm (Figure 1b).

Figure 1. Effect of stirring rate on (**a**) conversion of furfural and (**b**) 2-methylfuran yield over Ni/BC catalyst at reaction conditions. Initial concentration—0.39 M, tetralin solvent, temperature—240 °C, catalyst loading—0.3 g (particle size 106–250 μ m), H₂ pressure—50 bars, reaction time—5 h.

Nevertheless, the conversion of furfural was not significantly altered by further increase in stirring rate from 1000 rpm to 1200 rpm. Based on this, it appears that the application of a 1000 rpm stirring rate eliminates the external transport restriction in the solid–liquid–gas phase reaction. Consequently, this outcome is consistent with stirring rates that are frequently cited in the literature as being adequate to cause negligible external transport limitations in batch reactors and related reaction investigations [31–33]. As a result, data from future experiments were obtained with a mixing rate of 1000 rpm. This is because the mixing achieved by using a stirring rate of 1000 rpm can potentially achieve about 92% conversion and approximately 55% yield of 2-methylfuran, the product of interest in this study.

Internal mass transport constraints, such as pore diffusion or intraparticle diffusion, on the other hand, might influence the overall reaction rate. Hence, the synthesised Ni/BC catalyst powder was sieved into four different particle size ranges, such as >250 μ m, $212-250 \ \mu\text{m}$, $106-212 \ \mu\text{m}$, and $106 \ \mu\text{m}$ for the purpose of obtaining the particle size that offers negligible internal diffusion limitations. Based on sieved catalyst powder in the following particle size ranges, >250 μ m, 212–250 μ m, 106–212 μ m, and 106 μ m, the effect of intraparticle diffusion was studied. A similar approach was published in the literature [22,23,34]. Based on the conversion of furfural, the catalyst particle size range that results in the shortest diffusion length is determined. It is believed that the shorter the diffusion length to the active sites of the catalyst particles, the higher the conversion achieved, which indicates negligible intraparticle diffusion. The influence of catalyst particle size range on the conversion of furfural is shown in Figure 2. It can be observed that furfural conversion increases as the catalyst particle size decreases from greater than or equal to 250 µm to less than 106 µm. There is no doubt that catalyst particle size impacts the rate of furfural hydrogenation. The conversion approximately plateaus around 96% between 106–212 µm and <106 µm catalyst particles sizes. In fact, when reducing particle size from 106–212 μ m to less than 106 μ m, the conversion of furfural was not significantly affected. The result suggests that intraparticle diffusion or internal transport constraint are negligible when catalyst particles smaller than 106 μ m are used.





As the catalyst particle size decreases, the diffusion path length to the active sites also decreases. The resulting dispersed catalysts are made up of finely divided particles (less than 106 μ m), which gives them a very large surface area (353.5 m² g⁻¹), short diffusion length, and a high mass transfer rate for the catalytic hydrogenation of furfural into 2-methylfuran. Thus, under these conditions (i.e., 1000 rpm and catalyst particle size less 106 μ m), rapid hydrogen adsorption is encouraged by the presence of finely divided catalyst particles. This finding led to the usage of catalyst particles smaller than or equal to 106 μ m in subsequent studies for the gathering of experimental data for kinetic model development.

The conversion of furfural and the yields of products based on furfural hydrogenation over the Ni/BC catalyst with particle size less than or equal to 106 μ m and a stirring rate of 1000 rpm is shown in Figure 3. The identified products from the hydrogenation of furfural include furfuryl alcohol, 2-methyl furan, furan, and tetrahydrofuran. The result shows that the catalyst is more selective toward 2-methyl furan, which is the desired product with the highest yield. The reaction scheme can be summarised in Figure 4 with the identified products, 2-methylfuran, furfuryl alcohol, furan, and tetrahydrofuran. The results suggest that furfuryl alcohol is very quickly transformed into 2-methylfuran. As a result of the catalyst's hydrogenolysis abilities, furfural (C=O bond) was selectively hydrogenated, followed by cleavage and deoxygenation of the C–OH bond in furfuryl alcohol for the further production of 2-methylfuran. There is also the occurrence of the hydrogenolysis of C=O bonds, decarboxylation and hydrogenation of the C=O bond or the furan ring of furfural which resulted in the other observed products. It should be noted that the gaseous product was not analysed in this study.

The Ni/BC catalyst showed comparable conversion of furfural of about 92% to Mg-Al oxides supported by the Co-Ni bimetallic catalyst reported in the literature [35]. In comparison to the CuZnAl catalyst that produced a 70% yield of 2-methylfuran [11], the developed catalyst in this study produced a 60% yield. This result demonstrates the novel and efficient hydrogenation of furfural into 2-methylfurfuran using the developed Ni/BC catalysts.



Figure 3. Conversion and product yield of furfural hydrogenation over Ni/BC under the following reaction conditions: initial concentration—0.39 M, tetralin as solvent, temperature—230 °C, catalyst loading—0.3 g, particle size < 106 μ m, H₂ pressure—50 bars, stirring rate—1000 rpm, reaction time—4.5 h.



Figure 4. Reaction scheme for furfural hydrogenation over 15 wt% Ni/BC catalyst.

2.2. Effect of Catalyst Loading on Initial Reaction Rate

Based on the collected concentration data, a plot of concentration as a function of time was produced. The initial rates were estimated by a polynomial fitting approach, differentiated at time zero (t = 0). This approach has been used and reported in many studies [22,23,28,34,36]. Conversion into products was investigated as a function of catalyst loading. In Figure 5, the effect of catalyst loading on initial reaction rates of furfural hydrogenation is presented. It is observed from the graph that initial rate of reaction increases as the amount of catalyst increases. This is because as the amount of catalyst for the reaction increases, the number of active sites available for furfural and hydrogen adsorption and surface reaction increases likewise. Hence, a kinetically controlled regime should result in an increase in reaction rates in proportion to the number of catalytic active sites accessible [27].

It has been shown that both external and intraparticle diffusion transport constraints are negligible under reaction conditions of catalyst particle size below or equal to 106 μ m and stirring rate of 1000 rpm. To validate this finding, Figure 6 shows the Madon–Boudart test for furfural hydrogenation over a 15 wt% Ni/BC catalyst. According to the Madon–Boudart test, if reactions are carried out with internal transport restriction, the plot of ln r_0 vs. In (catalyst activity) will have a slope of 0.5, but if reactions are carried out with external transport limitation, it will have a slope of zero [27]. Catalyst activity in this case is

proportional to catalyst quantity, and external mass transfer is negligible. The reason for this is because the rate of the reaction is proportional to the number of catalytic active sites when mass transport restrictions are eliminated. This demonstrates a linear relationship between reaction $\ln r_0$ —ln (catalyst activity) due to Ni nanoparticle loadings on the catalyst surface. The correlation line of data plot for $\ln r_0$ vs. ln (catalyst activity) shown in Figure 6 has a slope of 0.64, which is greater than 0.5, which is an important indicator of internal mass transport limitations. Based on this finding, there is neither mass or heat transport resistance for a stirring rate of 1000 rpm and catalyst particle size of less than or equal to 106 µm.



Figure 5. Effect of catalyst loading on furfural hydrogenation reaction over Ni/BC at temperature 190 °C, stirring rate of 1000 rpm, hydrogen pressure of 50 bar, initial concentration of 0.39 M.



Figure 6. Madon–Boudart test for furfural hydrogenation reaction over Ni/BC at temperature 190 °C, stirring rate of 1000 rpm, hydrogen pressure of 50 bar, initial concentration of 0.39 M.

Under these conditions, the effect of catalyst loading on controlling resistance of gasliquid interfaces within batch reactors was also studied. The dissolution of hydrogen gas at the gas-liquid interface is described by Henry's law. Since the initial hydrogen pressure (i.e., 50 bar) was constant in all the experiments, the hydrogen concentration dissolved (C_H) was also constant. The technique of plotting (C_H/r_0) vs. 1/(catalyst loading) was utilised to determine the region of restriction in the gas–liquid interface within the system as detailed in the literature [37,38]. The slope signifies a combination of both external mass transfer and internal mass transport resistance at catalyst particles, while the intercept corresponds to the resistance to gas absorption at gas–liquid interfaces [38]. The result is presented in Figure 7. Resistance against gas absorption through the liquid film at the gas–liquid interface is represented by the intercept of the plot, which is 0.78 min, and the combined external mass transport restriction and internal pore diffusion restriction at the catalyst particle is 0.022 g min. Hence, the use of a more finely crushed catalyst particle decreases both internal and external mass transport resistances at catalyst particles at catalyst particle $\leq 106 \mu m$. This affirms the elimination of external and internal transport resistances under stirring rate 1000 rpm and $\leq 106 \mu m$; the resistance to absorption is insignificant, and surface reaction between furfural and hydrogen is most likely within the system constraints.



Figure 7. The plot of C_H/R_0 vs. 1/catalyst loading at a temperature of 200 °C, stirring rate of 1000 rpm, hydrogen pressure of 50 bar, initial concentration of 0.39 M.

2.3. Power Law Model Based on Initial Rate

Based on Figure 4, the formation of 2-methylfuran (MF) through furfural (FAL) hydrogenation via furfuryl alcohol (FOL) can be expressed as follows: $(FAL \rightarrow FOL \rightarrow 2MF)$ [11]. This suggests that furfuryl alcohol is produced right away from furfural and then converted into 2-methylfuran. The power law kinetic model is based on gathering data for furfural concentration as a function of time due to the intricacy of reaction pathways and difficulties in measuring the intermediate furfuryl alcohol during the hydrogenation process. The experimental data were collected at a constant hydrogen pressure of 50 bar. The effect of hydrogen pressure was not studied. Therefore, the rate of reaction terms of the concentration of furfural consumed per unit can be expressed using Equation (1):

$$r = \frac{dC}{dt} = kC^n \quad \text{or} \quad \ln r = \ln k + n\ln C, \tag{1}$$

where *r* denotes the reaction rate (M min⁻¹), *k* is the rate constant (M¹⁻ⁿ min⁻¹), *C* is the concentration of furfural (M) and *n* is the order of reaction.

The initial reaction rates (r_0) were computed using a polynomial fitting technique and then differentiated at time zero (t = 0) based on the plot of concentration as a function of time. To determine reaction rate constant (k) and order (n), the plot of natural log of the

initial rates $(\ln r_0)$ against natural log of initial furfural concentration $(\ln C_0)$ was produced for reaction temperatures 200 °C, 220 °C and 240 °C as shown in Figure 8. The order of reaction is the slope of the plot at each experimental temperature, while the corresponding rate constant is determined from the intercept.



Figure 8. Plots of $\ln r_0$ against $\ln C_0$ for reaction temperatures 200 °C, 220 °C, and 240 °C under the following reaction conditions: catalyst loading—0.2 g ($\leq 106 \mu$ m), hydrogen pressure—50 bar, tetralin as solvent, and stirring rate—1000 rpm.

The respective order of reaction (*n*) at reaction temperatures 200 °C, 220 °C and 240 °C is as follows: 0.42, 0.45 and 0.59, and the rate constants are 0.002695 $M^{0.58}$ min⁻¹, 0.003462 $M^{0.55}$ min⁻¹, and 0.005795 $M^{0.41}$ min⁻¹. The order of reaction is less than one, suggesting it is non-first order. The orders of reaction regarding furfural concentration vary from 0.42 to 0.6. Since the reaction cannot be described by a first-order reaction rate based on furfural initial concentration, the development of a model describing furfural hydrogenation reaction kinetics over the developed catalyst must consider processes occurring on the catalyst surface like adsorption, surface reaction, and desorption.

The Arrhenius equation, which is a thermodynamic function shown in Equation (2), was used to estimate activation energy. The plot of $\ln(k)$ versus 1/T to determine activation energy (E_a) and pre-exponential factor (A) is shown in Figure 9. The result shows that the rate constant increases as the reaction temperature increases from 200 °C to 260 °C. The estimated activation energy (E_a) is 40.22 kJ mol⁻¹ and the pre-exponential factor (A) is 70.26 min⁻¹,

$$\ln k = \ln A - \frac{E_a}{R} \left(\frac{1}{T}\right),\tag{2}$$

where *R* denotes the universal gas constant 8.314 J K⁻¹ mol⁻¹, E_a is the activation energy (J mol⁻¹), *A* is the pre-exponential factor (min⁻¹), and *k* is the rate constant (M^{0.55} min⁻¹).



Figure 9. The Arrhenius equation plot of ln(k) versus 1/T. Line shows a linear fit to the experimental data.

2.4. Langmuir-Hinshelwood Kinetic Type Model

The Langmuir–Hinshelwood (L–H) reaction rate model type equations are more adequate than power law to describe the reaction kinetics of furfural hydrogenation over the Ni/BC catalyst. This is because the L–H type rate model considers the active sites and processes such as adsorption, surface reaction, and desorption occurring on the surface of the catalyst that controls the reaction rate. However, previous studies have suggested a Langmuir-Hinshelwood-Hougen-Watson (LHHW) kinetic model for HDO or hydrogenation of bio-oil model compounds including furfural hydrogenation over an active metal supported catalyst that assumes either a single or a dual type of adsorption site and a surface reaction between a partially hydrogenated furfural species and an adsorbed hydrogen [23,25,34,36]. On the catalyst surface, it is assumed that on S_1 sites, atoms of H are the predominant species, while furfural dominates on S_2 sites. Based on this assumption, the reaction mechanism is illustrated by reaction schemes 1 to 4 for furfural conversion to 2-methylfuran. Typical L–H kinetic models can be invoked, which involves hydrogen dissociatively adsorbing on dual active sites and hydrogen chemisorbed on the same surface reacting with chemisorbed furfural. It has been reported that the most plausible fit is a dual-site mechanism with dissociative adsorption of hydrogen and surface reactions as rate-controlling steps [36], listed as follows:

- (i) molecular adsorption of furfural (F), furfuryl alcohol (FOL), and 2-methyl furan (2MF),
- (ii) dissociative adsorption of hydrogen on the catalyst active site,
- (iii) during the reaction, the partial pressure of hydrogen can be considered constant due to the excess of hydrogen,
- (iv) all adsorption sites are equivalent and independent of coverage, and
- (v) surface reaction is a rate-determining step.

It is believed that hydrogen adsorption onto the catalyst surface occurr dissociatively. Accordingly, Equations (3)–(6) represent the elementary steps taken in determining rate expressions for both competitive and non-competitive adsorption of dissociatively chemisorbed hydrogen (H_2) and furfural (F). Consequently, all adsorption and desorption sites of the catalysts are equivalent and independent of coverage [36].

$$H_2 + 2S_1 \stackrel{k_H}{\Leftrightarrow} 2H - S_1, \tag{3}$$

$$F + S_2 \stackrel{k_F}{\Leftrightarrow} F - S_2, \tag{4}$$

$$F-S_2 + H-S_1 \stackrel{\kappa_1}{\Leftrightarrow} S_2 - F.H-S_1, \tag{5}$$

$$S_2 - F.H - S_1 + H - S_1 \xrightarrow{\kappa_2} Product + S_2 + 2S_1.$$
(6)

The following rate expressions (Equations (7)–(9)) are derived for the selective hydrogenation of furfural (*F*) into 2-methylfuran based on dissociation of hydrogen.

Model-I: Competitive adsorption of dissociative H_2 and furfural (F) during hydrogenation:

$$r = \frac{k_s K_H K_F C_H C_F}{\left(1 + K_F C_F + K_H^{0.5} C_H^{0.5}\right)^3}.$$
(7)

Model-II: The model assumed one type of adsorption site (competitive adsorption) and the rate-determining step as a surface reaction between an adsorbed H and a partially hydrogenated furfural species:

$$r = \frac{k_s K_F K_H^{0.5} C_H^{0.5} C_F}{\left(1 + K_F C_F + K_H^{0.5} C_H^{0.5}\right)^2}.$$
(8)

Model-III: Non-competitive adsorption of dissociative H_2 and furfural during hydrogenation. Based on the assumption that *H* atoms dominate S_1 , and furfural dominates S_2 ,

$$r = \frac{k_s K_H K_F C_H C_F}{\left(1 + K_H^{0.5} C_H^{0.5}\right)^2 \left(1 + K_F C_F\right)},\tag{9}$$

where C_H denotes concentration of dissolved hydrogen in the solvent, C_F is the concentration of furfural (mol dm⁻³), k_s is the surface reaction rate constant (mol g⁻¹ min⁻¹), k_H is the hydrogen adsorption constant (m³ mol⁻¹), k_F is the furfural adsorption constant (m³ mol⁻¹), and r is the rate of reaction.

In the literature, the models of the two sites, Model-II and Model-III, have been applied to study furfural hydrogenation over copper dispersed on three forms of carbon, namely activated carbon, diamond and graphitised fibres [25]. In 2018, the hydrogenation of furfural derived from biomass into 2-methylfuran, which is a fuel additive, was studied over the Cu–Co/Al₂O₃ catalyst using Model-I and Model-II, respectively [36]. A linearisation of Equations (7)–(9) was conducted to determine the most appropriate model for the hydrogenation of furfural over Ni/BC catalysts. At constant hydrogen pressure of 50 bar (5 MPa), the concentration of dissolved hydrogen gas (C_H) can be determined with the aid of Henry's law ($C_H = K_H P_H$).

Therefore, the plot of C_{F0} vs. $\left(\frac{C_{F0}}{r_0}\right)^{\frac{1}{3}}$ (Model – I), C_{F0} vs. $\left(\frac{C_{F0}}{r_0}\right)^{\frac{1}{2}}$ (Model – II) and C_{F0} vs. $\frac{C_{F0}}{r_0}$ (Modell – III) should be a linear fit to the experimental data. Based on initial reaction rate obtained from the concentration time plots of the experimental data, Figure 10 shows the linear fits of the models to the data obtained at reaction temperatures 200 °C, 220 °C, and 240 °C.

The coefficient of determination (R^2) value, which represents the proportion of variation in the dependent variable that can be predicted completely from the independent variable, is used to evaluate the extent of linear fits to the initial reaction rate data. R^2 is a range of values between zero and one, and a value close to one implies that the dependent variable can be predicted efficiently based on the independent variable with less error expected. Therefore, based on the linearised fits to the initial rate data, the R^2 values are as follows: Model-I (0.990 (200 °C), 0.998 (220 °C) and 0.988 (240 °C)), Model-II (0.996 (200 °C), 0.999 (220 °C) and 0.995 (240 °C)), and Model-III (0.994 (200 °C), 0.999 (220 °C) and 0.995 (240 °C)). Model fitting and experimental data differ very little, possibly due to the narrow range of C_{F0}/r_0 . Based on R^2 values obtained through the linearised fits to the experimental data as shown in Figure 10, all three models predict the experimental data with certain degrees of error, but Model-III has higher R^2 values and minimised errors compared to Model-I and Model-II. This suggests that both Model-II and Model-III are more favourable kinetic models to describe the experimental data better than Model-I. Hence, the experimental data are studied using Model-III. It can be concluded that the best match for the experimental data is a dual-site mechanism with hydrogen dissociative adsorption and surface reaction as the rate-controlling step.



Figure 10. Fit of a linearised version of models to experimental data using initial rates obtained from their concentration time plots: (a) Model-I, (b) Model-II, and (c) Model-III (we note that C_0 is the same as C_{F0}).

2.5. Model Parameter Estimation

The sum of squared error (SSE) function was used for non-linear regression fit of model predictions (Model-III) by determining model parameters through the minimisation of residual sum of square (RSS) objective function shown in Equation (10). The model parameters were estimated by using the optimisation programme for a nonlinear generalised reduced gradient (GRG) solver in Microsoft Excel 2016. The RSS objective function was minimised to determine the model parameters that describe experimental data.

$$RSS = \sum_{i=1}^{n} (r_{Exp} - r_{Model})^2.$$
 (10)

By minimising the RSS objective error function using Microsoft Excel's GRG solver optimisation programme, kinetic Model-III parameters such as surface reaction rate constant (k_S) and adsorption constants $(K_H \text{ and } K_F)$ were determined, including heat of adsorption $(\Delta H_H \text{ and } \Delta H_F)$ and intrinsic activation energy (E_a). Table 2 shows the estimated kinetic model parameters for furfural hydrogenation reaction over the Ni/BC catalyst at a constant hydrogen pressure of 50 bar (5 MPa). With R^2 values beyond 95%, the proposed mechanistic model fit the data satisfactorily based on its kinetic model parameters (Table 2), as well as its correlation coefficients. In consequence, as reaction temperature increases, both rate constant and adsorption constant slightly increase. This indicates an increase in adsorption affinity and rate of reaction.

Table 2. Estimated values of kinetic model-III parameters for furfural hydrogenation reaction over the Ni/BC catalyst.

Temperature (°C)	k_S (mol g $^{-1}$ min $^{-1}$)	K_H (m ³ mol ⁻¹)	K_F (m ³ mol ⁻¹)	RSS
200	0.018	0.235	1.353	7.913×10^{-8}
220	0.023	0.254	1.388	8.876×10^{-8}
240	0.030	0.299	1.521	7.717×10^{-9}
260	0.036	0.320	1.582	$3.430 imes 10^{-8}$

Note: RSS—residual sum of squares, k_s —surface reaction rate constant (mol g⁻¹ min⁻¹), k_H —hydrogen adsorption constant (m³ mol⁻¹), and k_F —furfural adsorption constant (m³ mol⁻¹).

Figure 11 shows the plot of experimental reaction rates against model predicted rates for Model-III, which indicates a strong agreement between experimental reaction rates and predicted reaction rates. This claim can be supported by a correlation coefficient greater than 99% between the experimental rates and the predicted rates from the model. Hence, a good correlation is found between the predicted values by the model and experimental results. It can be concluded that the experimental results could also be mechanistically explained using Langmuir–Hinshelwood type kinetic models.



Figure 11. Parity plot of Model-III to compare predicted rates by model and experimentally determined rates. Line shows linear fit to experimental data points.

The activation energy for the hydrogenation reaction of furfural was determined using Arrhenius Equation (2), which relates reaction rate constant with temperature. Based on linearised Arrhenius equation, the plot of $\ln k_S$ vs. 1/T for reaction temperatures from 200 °C to 260 °C is shown in Figure 12. However, the Van't Hoff isochore expression (Equation (11)) was used to estimate the enthalpy of adsorption as shown in Figure 13a,b for both furfural and hydrogen.

Van't Hoff isochore equation is as follows:

$$\ln K_{ads} = \left(\frac{-\Delta H_{ads}}{RT}\right) + \left(\frac{\Delta S}{R}\right),\tag{11}$$

where *R* denotes the universal gas constant 8.314 J K⁻¹ mol⁻¹, E_a is activation energy (J mol⁻¹), *A* is the pre-exponential factor, K_{ads} is the adsorption equilibrium constant, ΔH_{ads} is the enthalpy changes due to adsorption, and ΔS is the change in entropy of the system.



Figure 12. Arrhenius plot for furfural hydrogenation reaction over the Ni/BC catalyst.



Figure 13. Van't Hoff isochore plots: (**a**) furfural adsorption onto the catalyst, and (**b**) hydrogen adsorption onto the Ni/BC catalyst.

Based on the Arrhenius plot, the estimated activation energy is 24.39 kJ mol⁻¹. It has been found that the apparent energy of activation for the surface reaction is 11.8 kcal mol⁻¹ (49.4 kJ mol⁻¹) for furfural hydrogenation reaction over the Cu/SiO₂ catalyst [39]. The active metals and support material properties can account for the differences. However, activation energy estimates indicate that experimental data were collected in a kinetically controlled regime, because they are lower than typical activation energies. Thus, due to the low activation energy of the reaction over the Ni/BC catalyst, it implies that the reaction proceeds quickly as many reactants possess the required energy. In other words, the Ni/BC catalyst is a material that is added to lower the reaction's activation energy to speed up the reaction. Catalysis occur since the activation energy is reduced, thus speeding up the reaction. This confirms the conclusion that external and internal mass transport are negligible under the experimental conditions. On the other hand, the enthalpy of adsorption and entropy change for the adsorption of furfural on the active sites of the Ni/BC catalyst are -5.86 kJ mol⁻¹ and 14.81 J K⁻¹ mol⁻¹, respectively, while the values of adsorption of hydrogen enthalpy and entropy change are -11.43 kJ mol⁻¹ and 12.04 J K⁻¹ mol⁻¹. It can be seen from the plots that furfural adsorbs on Ni/BC with a higher heat of adsorption than hydrogen. Compared with hydrogen, furfural shows a stronger adsorption on the catalyst surface. The enthalpy of adsorption shows that heat is released during physisorption adsorption of hydrogen and furfural to the surface of the biochar. Thermodynamically, adsorption is more favourable since it is an exothermic process [40]. It agrees with the exothermic process involved in adsorption, and similar values have been reported for hydrogen and furfural adsorption [35]. Notwithstanding, the experimental results obtained for the reaction indicate that at a higher temperature, all components have greater adsorption constant values, which is consistent with the observation (Table 2).

2.6. Catalyst Reusability

Reusability of catalysts is essential for improving process economics. The reusability of the Ni/BC catalyst was studied by recovering the spent catalyst and reusing the catalysts for four successive furfural hydrogenation reaction cycles. The recovered catalysts were dried at 200 °C for 30 min to remove unreacted furfural, solvent, and products from the catalyst pores, before being used in subsequent experiment. The reusability of the Ni/BC catalyst was studied over four successive cycle reactions at a temperature of 230 °C, initial concentration of 0.39 M, hydrogen gas pressure of 50 bar, catalyst loading pf 0.3 g, stirring rate of 1000 rpm and reaction time of 5 h. Figure 14 displays changes in conversion and product yields for each reaction cycle. As can be seen from the graph, there is a slight noticeable difference between the first and fourth cycles when it comes to furfural conversion (99% to 90%) and the yield of 2-methyl furan (57% to 54%). the yields of furan and tetrahydrofuran follow the same trend. The catalyst is stable as evidenced by this observation. On the other hand, the yield of furfuryl alcohol increases as the reaction cycle increases from first to fourth cycle. Based on XRD patterns for the catalyst spent in the first cycle experiment reported in a previous publication [21], the peak positions and intensities of Ni/BC catalysts did not change, which indicates that over the course of reaction time, no phase transformation or deactivation occurred due to furfural hydrogenation. For more insight into the reason for gradual decrease in furfural conversion and 2-methylfuran yield, it is worthwhile to systematically characterise the spent after each reaction cycle.



Figure 14. Effect of catalyst reusability on conversion and product yield from furfural hydrogenation over Ni/BC at the following reaction conditions: initial concentration—0.39 M, tetralin as solvent, temperature—230 °C, catalyst loading—0.3 g, stirring rate—1000 rpm, H₂ pressure—50 bars, reaction time—5 h.

Regarding the observed decline in furfural conversion and the yield of 2-methyl furan, this occurs possibly due to the reduced strength in the catalyst active sites after successive reaction and drying steps [32]. This is further reaffirmed by the increasing yield of furfuryl

alcohol from first cycle to fourth cycle, signifying that the hydrogenation of furfuryl alcohol to 2-methyl furan is impacted. This can be attributed to the chemisorbed aromatic species (furfural, furan, 2-methyl furan, etc.) on the catalyst active sites after consecutive reaction, which could have adversely impacted the strength since no appropriate regeneration of catalysts was performed before reuse. Additionally, it has been proposed that aromatic molecules that are adsorbed onto the catalyst's active sites during HDO or the hydrogenation of bio-oil oxygenate model compounds may result in a decrease in the catalyst's activity in subsequent reaction cycles [41]. This explains the gradual decrease in the yield of 2-methyl furan and, correspondingly, in the yield of furfuryl alcohol. In contrast, a solvent such as acetone can be used to wash the spent catalyst before drying at 200 °C in order to remove most of the bound aromatics and furans.

2.7. Effect of Acidification Time

In a prior study published in our previous work, it was found that treating the BC with sulfuric acid (H_2SO_4) greatly improved the surface area, pore volume, and mesoporosity [21]. Therefore, acid treatment of a catalyst could increase reaction rates by improving mass transport and providing more surface area for adsorption and surface reactions. The surface area and porosity of catalysts are believed to increase with increasing acid time (Table 1). Figure 15 demonstrates how BC duration in the sulfuric acid affects the catalytic performance of the Ni/BC catalyst in terms of the yield of products and furfural conversion. Based on furfural conversion and 2-methyl furan yield, the acid-treated Ni/BC catalyst outperforms the non-treated (NT) counterpart. This is consistent with the improvement in the physicochemical characteristics of the catalyst observed in prior published work after chemical treatment of the BC [21,42,43]. It has been observed that a sulfuric-activated BC catalyst has excellent mesopores and a high surface area, thereby improving the adsorptive effect and therefore catalytic activity.



Figure 15. Conversion and product yield of furfural hydrogenation over Ni/BC at the following reaction conditions: initial concentration—0.39 M, tetralin as solvent, temperature—240 °C, catalyst loading—0.3 g, stirring rate—1000 rpm, H₂ pressure—50 bars, and reaction time—5 h. (we note: NT denotes no acid treatment, and 3 h, 6 h, and 9 h are 3, 6 and 9 h acid treatments).

Notably, when the sulfuric acid treatment duration of the BC employed in the catalyst development increased from 3 to 9 h, the conversion of furfural and the yield of 2-methyl furan increased slightly. An increase in the yield of furan and tetrahydrofuran can also be observed, along with a decrease in the yield of furfuryl alcohol. In 9 h sulfuric acid treatment of BC, approximately 100% of furfural is converted, and 71% of 2-methyl furan

is yielded. Plausible explanation can be attributed to increased mesoporosity of the BC support following acid treatment [21], resulting in the enhancement of mass transport, activity, and selectivity for 2-methyl furan (Figure 15). This is supported by the fact that the yield of furfuryl alcohol, a required product for producing 2-methyl furan, diminishes with increasing acid treatment duration. The improvement in catalyst pore size following 9 h of acid treatment can also be favourable for high-molecular-weight hydrocarbon tetralin used as a solvent due to the enhanced intraparticle diffusion. A greater mass transport efficiency favours the kinetic control regime, benefiting adsorption and surface reactions, respectively. It appears that the reaction resulting in pore creation with sulfuric acid of biochar is slow, since sulfuric acid activation time shows significant change in furfural conversion and yield of products.

3. Materials and Methods

3.1. Materials

The following are details on the purity and suppliers of chemical compounds, materials, and gases used in this study: nickel (II) nitrate hexahydrate (Ni(NO₃)₂·6H₂O, 99.99%), furfural (99%), 2-methylfuran (99%), furfuryl alcohol (98%), furan (99%), and tetrahydrofuran (99%) were purchased from Sigma Aldrich, Gillingham, UK, tetralin (>98%) and sulphuric acid, H₂SO₄ (>95%) were purchased from Fisher Scientific, Loughborough, UK, and bottled hydrogen gas and nitrogen gas were purchased from British Oxygen Company (BOC), Ltd., Woking, UK. The biochar (BC) employed as a catalyst support in this study was synthesised at the Hohenburg plant in Germany through the pyrolysis of sewage sludge waste.

3.2. BC Modification

About 20 g of the BC was sulfonated in a 250 mL round-bottomed flask fitted with a reflux condenser at 120 °C on a hot plate equipped with a thermocouple and a heating mantle for 3 h with 300 mL concentrated sulfuric acid. The precipitate was repeatedly washed with hot distilled water until sulphate ions were removed from the filtrate, as confirmed by a pH of 7. The mixture was filtered using a vacuum filter when it cooled to room temperature. The samples were oven dried overnight at 80 °C, and the solid substance generated was labelled as BC. A full description of BC sulfuric acid activatedbiochar has been published elsewhere [21]. As a result of the reaction between the acid and oxygenated molecules in the BC, this treatment improves the surface of the BC by constructing new pores while expanding micropores.

3.3. Catalyst Synthesis

An appropriate amount of nickel (II) nitrate hexahydrate was dissolved in 50 mL of deionised water, and 20 g of sulfuric acid-activated BC was added to prepare a 15 wt% Ni/BC catalyst. The mixture was stirred with a magnet stirrer at 800 rpm and 70 °C for 1 h when all surface water evaporated, leaving the wet solid catalyst, which was then dried overnight at 80 °C. To develop a Ni/BC catalyst, the resultant material was subsequently calcined at 500 °C for 5.5 h. The catalyst was pre-activated ex situ using a Carbolite Geo TF1200 tubular furnace, Carbolite Gero, Hope Valley, UK, for 5 h at 500 °C with a steady flow of 5 vol.% H_2/N_2 .

3.4. Catalyst Characterisation

A Micromeritrics Analytical Instrument ASAP 2010, Micromeritics, Norcross GA, USA was used to measure the developed Ni/BC catalyst's surface area, pore volume, and pore size distribution through the nitrogen adsorption–desorption technique. A full description of the catalyst characterisation techniques used in this study has been published elsewhere [21]. Based on the available sieve size, the catalyst powder was sieved into the following particle size ranges: >250 μ m, 212–250 μ m, 106–212 μ m and <106 μ m, respectively.

3.5. Catalytic Reaction and Product Quantification

The HDO reaction was carried out under the following range of conditions: temperatures of 100 °C to 150 °C, hydrogen (H₂) pressures of 30 to 50 bar and stirring rate of 1000 rpm. Based on preliminary experiments, it was found that a stirring rate of 1000 rpm was sufficient in eliminating the effect of external mass transfer. Hence, all experiments were performed using this mixing speed. As soon as hydrogen was introduced into the reactor, reaction time was started. For each reaction, samples were taken at the end (undisturbed) or in equally spaced time intervals throughout the duration of the reaction, and the amount of each sample was fixed to reduce pressure loss. For furfural hydrogenation experiments, this study employed a 100 mL stainless steel batch reactor (the Parr reactor 4598 system, Parr, Moline IL, USA) which was provided by Parr Instruments. The reactor was fed with an appropriate amount of the developed catalyst (0–0.4 g), an initial furfural concentration of 0.39 M in 40 mL of the solvent (tetralin), sealed, and purged three times with nitrogen before being pressurised to a 50 bar pressure with hydrogen gas. Due to its ability to dissolve furfural and its hydrogenation products as well as its thermal stability within the temperature range examined, tetralin was selected as the solvent. Additionally, it has a gas venting valve and a liquid sampling valve for collecting samples during experiments and purging gases at the start and end of experiments. The experiment was carried out at the following reaction temperatures: 200 °C, 220 °C, and 240 °C, respectively, at a stirring rate of 1000 rpm. Prior to recycling the spent catalyst when carrying out the catalyst reusability investigation, after each of the four subsequent furfural hydrogenation experiment cycles, the recovered catalyst was dried at 200 °C for 30 min in order to remove unreacted furfural, solvent, and products from the pores, before being for experiment without further treatment. In some experiments, samples were collected at the end, whereas in others, samples were collected during time-on-process at time intervals for the purpose of concentration—time plots and initial reaction rate. Approximately 2 mL of liquid samples were collected at regular intervals from the reactor for each experiment and analysed using a gas chromatographic (GC) technique (Agilent GC 6890N, Agilent, Santa Clara, USA) equipped with flame ionisation detector (FID) and a Zebron ZB-Wax capillary column $(30 \text{ m} \times 0.320 \text{ mm a} \times 0.25 \text{ }\mu\text{m})$, Phenomenex, Torrance, CA, USA. The injector and detector temperatures were set at 320 °C and 330 °C, respectively, with helium serving as the carrier gas. Each sample analysis was performed with 1 μ L of the collected sample injected into the column using a split ratio of 20:1. A full description of the GC-FID analysis method has been reported elsewhere [21]. Calibration curves based on the chemical compounds of typical products such as furfural, 2-methylfuran, furfuryl alcohol, furan, and tetrahydrofuran anticipated from furfural hydrogenation prior to sample GC analysis were developed for identification and quantification. Quantification of the conversion of furfural, the yields of products, and their selectivity was performed using the following equations:

$$Conversion (\%) = \frac{\text{initial moles of reactant } (n_{r0}) - \text{final moles of reactant } (n_{r1})}{\text{initial moles of reactnt } (n_{r0})} \times 100,$$
(12)

Product yield (%) =
$$\frac{\text{Moles of desired product } (n_p)}{\text{initial moles of reactant } (n_{r0})} \times 100,$$
 (13)

Selectivity (%) =
$$\frac{\text{Moles of desired product } (n_p)}{\text{total moles of products}} \times 100.$$
 (14)

4. Conclusions

The current study used a chemical activation method to modify BC textural characteristics to achieve a greatly enhanced specific surface area, mesopore structure, and pore volume in comparison to its pristine BC. In this study, sulfuric acid modified BC was used to develop a Ni/BC catalyst that can selectively hydrogenate furfural into 2-methylfuran. In comparison with previous catalysts, Ni/BC is a cost-effective catalyst with comparative performance that enables circular economy, sustainability, and industrial scale application in bio-oil catalytic upgrading. Using a stirring rate of 1000 rpm and a catalyst particle size of less than or equal to 106 μ m, both external and intra-particle diffusion became insignificant, ensuring that the experiments were carried out under a kinetic controlled regime.

Three plausible Langmuir–Hinshelwood–Hougen–Watsson (LHHW) kinetic type models, competitive (Model-I), with a competitive adsorption site (Model-II) and a noncompetitive adsorption site (Model-III), were screened based on the fact that an R^2 value greater than 99% demonstrated that the experimental data were satisfactorily fit to the model. It was shown that the best fit is Model-III, which is a dual-site adsorption mechanism involving furfural adsorption as well as hydrogen dissociative adsorption and surface reactions. Based on the results of four successive experimental studies, the developed Ni/BC catalyst demonstrated excellent stability in terms of furfural conversion (96%) and 2-methylfuran yield (54%) at the fourth experiment compared to 99% and 57% for the first experiment. To gain further insight into the reason for the gradual decline in Ni/BC catalyst performance in terms of furfural conversion and 2-methylfuran yield, it would be beneficial to characterise the spent catalyst after each reaction cycle systematically.

Supplementary Materials: The following supporting information can be downloaded at: https://www.mdpi.com/article/10.3390/catal14010054/s1, Table S1: Surface area, pore volume and Ni nanoparticle content. Figure S1: TEM analysis and histograms of the Ni metal nanoparticle sizes of Ni/BC. Figure S2: FT-IR spectra of biochar and biochar (H₂SO₄), and their corresponding Ni catalysts. Figure S3: CO₂-TPD profile of raw Ni/biochar and sulfuric acid activated biochar Ni/BC catalysts.

Author Contributions: Conceptualisation, J.W.; methodology, I.M.; software, I.M.; validation, I.M. and J.W.; formal analysis, I.M.; investigation, I.M.; resources, J.W.; data curation, I.M.; writing—original draft preparation, I.M.; writing—review and editing, I.M., A.H., A.I. and J.W.; visualisation, I.M.; supervision, A.I. and J.W.; project administration, J.W.; funding acquisition, I.M. and J.W. All authors have read and agreed to the published version of the manuscript.

Funding: Financial support for this work was provided by Petroleum Technology Development Fund (PTDF), Nigeria and School of Chemical Engineering, University of Birmingham.

Data Availability Statement: Data are available free of charge via https://edata.bham.ac.uk.

Acknowledgments: Technical support from Jonathan Goldfinch and ChyntolKanhimbe of School of Chemical Engineering, University of Birmingham, John Wedderburn (Metallurgy and Materials), University of Birmingham and June Callison, Lisa Allen and Sining Chen (Harwell Research complex) is well acknowledged.

Conflicts of Interest: The authors declare no conflict of interest.

References

- 1. Dong, F.; Zhu, Y.; Zheng, H.; Zhu, Y.; Li, X.; Li, Y. Cr-free Cu-catalysts for the selective hydrogenation of biomass-derived furfural to 2-methylfuran: The synergistic effect of metal and acid sites. *J. Mol. Catal. A Chem.* **2015**, *398*, 140–148. [CrossRef]
- Durndell, L.J.; Zou, G.; Shangguan, W.; Lee, A.F.; Wilson, K. Structure-Reactivity Relations in Ruthenium Catalysed Furfural Hydrogenation. *ChemCatChem* 2019, 11, 3927–3932. [CrossRef]
- Li, C.; Xu, G.; Liu, X.; Zhang, Y.; Fu, Y. Hydrogenation of biomass-derived furfural to tetrahydrofurfuryl alcohol over hydroxyapatite-supported Pd catalyst under mild conditions. *Ind. Eng. Chem. Res.* 2017, 56, 8843–8849. [CrossRef]
- O'Driscoll, A.i.; Curtin, T.; Hernández, W.Y.; Van Der Voort, P.; Leahy, J.J. Hydrogenation of furfural with a Pt–Sn catalyst: The suitability to sustainable industrial application. Org. Process Res. Dev. 2016, 20, 1917–1929. [CrossRef]
- Wang, Y.; Zhao, D.; Rodríguez-Padrón, D.; Len, C. Recent advances in catalytic hydrogenation of furfural. *Catalysts* 2019, 9, 796. [CrossRef]
- Lee, W.-S.; Wang, Z.; Zheng, W.; Vlachos, D.G.; Bhan, A. Vapor phase hydrodeoxygenation of furfural to 2-methylfuran on molybdenum carbide catalysts. *Catal. Sci. Technol.* 2014, *4*, 2340–2352. [CrossRef]
- 7. Yang, Y.; Chen, L.; Chen, Y.; Liu, W.; Feng, H.; Wang, B.; Zhang, X.; Wei, M. The selective hydrogenation of furfural over intermetallic compounds with outstanding catalytic performance. *Green Chem.* **2019**, *21*, 5352–5362. [CrossRef]
- 8. Date, N.S.; Hengne, A.M.; Huang, K.-W.; Chikate, R.C.; Rode, C.V. Single pot selective hydrogenation of furfural to 2-methylfuran over carbon supported iridium catalysts. *Green Chem.* **2018**, *20*, 2027–2037. [CrossRef]
- 9. Chen, X.; Zhang, L.; Zhang, B.; Guo, X.; Mu, X. Highly selective hydrogenation of furfural to furfuryl alcohol over Pt nanoparticles supported on g-C₃N₄ nanosheets catalysts in water. *Sci. Rep.* **2016**, *6*, 28558. [CrossRef]

- 10. Srivastava, S.; Jadeja, G.; Parikh, J. A versatile bi-metallic copper-cobalt catalyst for liquid phase hydrogenation of furfural to 2-methylfuran. *RSC Adv.* **2016**, *6*, 1649–1658. [CrossRef]
- 11. Niu, H.; Luo, J.; Li, C.; Wang, B.; Liang, C. Transfer hydrogenation of biomass-derived furfural to 2-methylfuran over CuZnAl catalysts. *Ind. Eng. Chem. Res.* 2019, *58*, 6298–6308. [CrossRef]
- 12. Chang, X.; Liu, A.F.; Cai, B.; Luo, J.Y.; Pan, H.; Huang, Y.B. Catalytic transfer hydrogenation of furfural to 2-methylfuran and 2-methyltetrahydrofuran over bimetallic copper–palladium catalysts. *ChemSusChem* **2016**, *9*, 3330–3337. [CrossRef]
- 13. Li, S.; Fan, Y.; Wu, C.; Zhuang, C.; Wang, Y.; Li, X.; Zhao, J.; Zheng, Z. Selective hydrogenation of furfural over the Co-based catalyst: A subtle synergy with Ni and Zn dopants. *ACS Appl. Mater. Interfaces* **2021**, *13*, 8507–8517. [CrossRef] [PubMed]
- 14. Taylor, M.J.; Durndell, L.J.; Isaacs, M.A.; Parlett, C.M.; Wilson, K.; Lee, A.F.; Kyriakou, G. Highly selective hydrogenation of furfural over supported Pt nanoparticles under mild conditions. *Appl. Catal. B Environ.* **2016**, *180*, 580–585. [CrossRef]
- 15. Huš, M.; Likozar, B.; Grilc, M. Furfural hydrogenation over Cu, Ni, Pd, Pt, Re, Rh and Ru catalysts: Ab initio modelling of adsorption, desorption and reaction micro-kinetics. *Chem. Eng. J.* 2022, 436, 135070.
- 16. Yan, K.; Wu, G.; Lafleur, T.; Jarvis, C. Production, properties and catalytic hydrogenation of furfural to fuel additives and value-added chemicals. *Renew. Sustain. Energy Rev.* **2014**, *38*, 663–676. [CrossRef]
- 17. Jahangiri, H.; Bennett, J.; Mahjoubi, P.; Wilson, K.; Gu, S. A review of advanced catalyst development for Fischer–Tropsch synthesis of hydrocarbons from biomass derived syn-gas. *Catal. Sci. Technol.* **2014**, *4*, 2210–2229. [CrossRef]
- 18. Quan, C.; Zhou, Y.; Wang, J.; Wu, C.; Gao, N. Biomass-based carbon materials for CO₂ capture: A review. J. CO2 Util. 2023, 68, 102373. [CrossRef]
- 19. Ren, S.; Lei, H.; Wang, L.; Bu, Q.; Chen, S.; Wu, J. Hydrocarbon and hydrogen-rich syngas production by biomass catalytic pyrolysis and bio-oil upgrading over biochar catalysts. *RSC Adv.* **2014**, *4*, 10731–10737. [CrossRef]
- Wu, P.; Singh, B.P.; Wang, H.; Jia, Z.; Wang, Y.; Chen, W. Bibliometric analysis of biochar research in 2021: A critical review for development, hotspots and trend directions. *Biochar* 2023, 5, 6. [CrossRef]
- 21. Mudi, I.; Hart, A.; Ingram, A.; Wood, J. Catalytic hydrodeoxygenation of vanillin, a bio-oil model compound over renewable Ni/biochar catalysts. *Catalysts* **2023**, *13*, 171. [CrossRef]
- Aliu, E.; Hart, A.; Wood, J. Kinetics of vanillin hydrodeoxygenation reaction in an organic solvent using a Pd/C catalyst. *Ind. Eng. Chem. Res.* 2019, 58, 15162–15172. [CrossRef]
- 23. Maccarrone, M.J.; Torres, G.; Lederhos, C.; Betti, C.; Badano, J.M.; Quiroga, M.; Yori, J. Kinetic study of the partial hydrogenation of 1-heptyne over Ni and Pd supported on alumina. *Hydrogenation* **2012**, 159–184. [CrossRef]
- 24. Pirmoradi, M.; Kastner, J.R. A kinetic model of multi-step furfural hydrogenation over a Pd-TiO₂ supported activated carbon catalyst. *Chem. Eng. J.* **2021**, *414*, 128693. [CrossRef]
- Rao, R.S.; Baker, R.T.K.; Vannice, M.A. Furfural hydrogenation over carbon-supported copper. *Catal. Lett.* 1999, 60, 51–57. [CrossRef]
- 26. Stamatiou, I.K.; Muller, F.L. Determination of mass transfer resistances of fast reactions in three-phase mechanically agitated slurry reactors. *AIChE J.* **2017**, *63*, 273–282. [CrossRef]
- Singh, U.K.; Vannice, M.A. Kinetics of liquid-phase hydrogenation reactions over supported metal catalysts—A review. *Appl. Catal. A Gen.* 2001, 213, 1–24. [CrossRef]
- Aliu, E. Vanillin Hydrodeoxygenation Reaction Kinetics over Supported Monometallic and Bimetallic Pd-Based Catalysts in Different Solvents and Binary Environments. Ph.D. Thesis, University of Birmingham, Birmingham, UK, 2020.
- 29. Quitian, A.; Ancheyta, J. Experimental methods for developing kinetic models for hydrocracking reactions with slurry-phase catalyst using batch reactors. *Energy Fuels* **2016**, *30*, 4419–4437. [CrossRef]
- Chen, Y.; Miller, D.J.; Jackson, J.E. Kinetics of aqueous-phase hydrogenation of organic acids and their mixtures over carbon supported ruthenium catalyst. *Ind. Eng. Chem. Res.* 2007, 46, 3334–3340. [CrossRef]
- Aliu, E.; Hart, A.; Wood, J. Mild-Temperature hydrodeoxygenation of vanillin a typical bio-oil model compound to Creosol a potential future biofuel. *Catal. Today* 2021, 379, 70–79. [CrossRef]
- 32. Lawal, A.M.; Hart, A.; Daly, H.; Hardacre, C.; Wood, J. Catalytic hydrogenation of short chain carboxylic acids typical of model compound found in bio-oils. *Ind. Eng. Chem. Res.* 2019, *58*, 7998–8008. [CrossRef]
- Lawal, A.M.; Hart, A.; Daly, H.; Hardacre, C.; Wood, J. Kinetics of hydrogenation of acetic acid over supported platinum catalyst. Energy Fuels 2019, 33, 5551–5560. [CrossRef]
- Aliu, E.; Hart, A.; Wood, J. Reaction kinetics of vanillin hydrodeoxygenation in acidic and nonacidic environments using bimetallic PdRh/Al₂O₃ catalyst. *Energy Fuels* 2019, 33, 11712–11723. [CrossRef]
- 35. Gong, X.; Li, N.; Li, Y.; Hu, R. The catalytic hydrogenation of furfural to 2-methylfuran over the Mg-Al oxides supported Co-Ni bimetallic catalysts. *Mol. Catal.* **2022**, *531*, 112651. [CrossRef]
- 36. Srivastava, S.; Jadeja, G.; Parikh, J. Copper-cobalt catalyzed liquid phase hydrogenation of furfural to 2-methylfuran: An optimization, kinetics and reaction mechanism study. *Chem. Eng. Res. Des.* **2018**, *132*, 313–324. [CrossRef]
- 37. Winterbottom, J.M.; King, M. Reactor Design for Chemical Engineers; CRC Press: Boca Raton, FL, USA, 1999.
- 38. Joseph, W. Three-phase catalytic reactors for hydrogenation and oxidation reactions. Phys. Sci. Rev. 2016, 1, 20150019. [CrossRef]
- 39. Sitthisa, S.; Sooknoi, T.; Ma, Y.; Balbuena, P.B.; Resasco, D.E. Kinetics and mechanism of hydrogenation of furfural on Cu/SiO₂ catalysts. *J. Catal.* **2011**, 277, 1–13. [CrossRef]

- 40. Torres-Knoop, A.; Poursaeidesfahani, A.; Vlugt, T.J.; Dubbeldam, D. Behavior of the enthalpy of adsorption in nanoporous materials close to saturation conditions. *J. Chem. Theory Comput.* **2017**, *13*, 3326–3339. [CrossRef]
- 41. Pipitone, G.; Zoppi, G.; Bocchini, S.; Rizzo, A.M.; Chiaramonti, D.; Pirone, R.; Bensaid, S. Aqueous phase reforming of the residual waters derived from lignin-rich hydrothermal liquefaction: Investigation of representative organic compounds and actual biorefinery streams. *Catal. Today* **2020**, *345*, 237–250. [CrossRef]
- 42. Ahiduzzaman, M.; Sadrul Islam, A. Preparation of porous bio-char and activated carbon from rice husk by leaching ash and chemical activation. *SpringerPlus* **2016**, *5*, 1–14. [CrossRef]
- 43. Iriarte-Velasco, U.; Sierra, I.; Zudaire, L.; Ayastuy, J.L. Preparation of a porous biochar from the acid activation of pork bones. *Food Bioprod. Process.* **2016**, *98*, 341–353. [CrossRef]

Disclaimer/Publisher's Note: The statements, opinions and data contained in all publications are solely those of the individual author(s) and contributor(s) and not of MDPI and/or the editor(s). MDPI and/or the editor(s) disclaim responsibility for any injury to people or property resulting from any ideas, methods, instructions or products referred to in the content.