Rapid Microwave-Assisted Polyol Synthesis of TiO₂-Supported Ruthenium Catalysts for Levulinic Acid Hydrogenation

Alexander G. R. Howe, Rhodri Maunder, David J. Morgan and Jennifer K. Edwards *

School of Chemistry, Cardiff University, Main Building, Park Place, Cardiff CF10 3AT, UK
* Correspondence: EdwardsJK@cardiff.ac.uk

Received: 7 August 2019; Accepted: 28 August 2019; Published: 5 September 2019

Abstract: One wt% Ru/TiO₂ catalysts prepared by a one-pot microwave-assisted polyol method have been shown to be highly active for Levulinic acid hydrogenation to γ-Valerolactone. Preparation temperature, microwave irradiation time and choice of Ru precursor were found to have a significant effect on catalyst activity. In the case of Ru(acac)₃-derived catalysts, increasing temperature and longer irradiation times increased catalyst activity to a maximum LA conversion of 69%. Conversely, for catalysts prepared using RuCl₃, shorter preparation times and lower temperatures yielded more active catalysts, with a maximum LA conversion of 67%. Catalysts prepared using either precursor were found to contain highly dispersed nanoparticles <3 nm in diameter. XPS analysis of the most and least active catalysts shows that the catalyst surface is covered in a layer of insoluble carbon with surface concentrations exceeding 40% in some cases. This can be attributed to the formation of large condensation oligomers from the reaction between the solvent, ethylene glycol and its oxidation products, as evidenced by the presence of C-O and C=O functionality on the catalyst surface.

Keywords: levulinic acid hydrogenation; microwave; ruthenium catalyst

1. Introduction

Biomass-derived feedstocks currently show potential as environmentally benign alternatives to fossil fuel derived chemicals, either directly as a fuel or as an intermediate for incorporation into existing commodity scale chemical processes [1]. The conversion of bulk biomass into useful chemicals has been extensively explored over the past decade, with an emphasis on utilization of lignocellulosic biomass [2]. Cellulose, a major (35–50%) component of such biomass, is readily hydrolyzed under acidic conditions to Levulinic acid (LA). LA was identified by the U.S. Department of Energy in 2004 as a bio-derived platform chemical capable of incorporation into traditionally petroleum derived value chains [3]. The application of Levulinic acid derived species and the prerequisite chemical transformations have been thoroughly explored [4]. One such transformation is the hydrogenation of LA to γ-Valerolactone (GVL), schematically presented in Figure 1, with GVL itself also being considered a viable platform molecule for the production of fuels or fuel additives [5].
The production of GVL from LA has been long known to occur readily in both the gas and liquid phase, the former having been investigated by the Quaker Oats Company as early as the 1950s, and the latter having been reported to occur over PtO₂ in 1930 [6,7]. Over the subsequent decades, further work sought to develop catalysts which retained high activity at lower temperatures and pressures, initially using bulk materials, such as Raney Nickel and later on the application of nanomaterials [8,9]. More recently, supported precious metal catalysts have been exhaustively explored for levulinic acid hydrogenation, showing promising potential for commercialization of the process. Early work by Manzer et al. found that carbon-supported Ir, Rh, Pt, Pd, Re, Ni and Ru nanoparticle catalysts were active for LA hydrogenation. Supported Pt, Re and Ni catalysts were found to be relatively inactive, exhibiting conversions below 20%, whereas the analogous Ir, Rh and Pd catalysts displayed conversions up to 40%. The 5 wt% Ru/C in particular exhibited both a LA conversion of 80% and a 91% selectivity towards GVL [10]. At the same time, Yan and co-workers compared the activity of Ru/C and Pd/C catalysts to Raney and Urushibara nickel, showing that both the Pd/C and bulk nickel catalysts were considerably less selective towards GVL than the Ru/C analogue [11].

Later work by Upare and co-workers showed that the poor selectivity of supported Pt/C and Pd/C catalysts was due to their relative inactivity towards the hydrogenation of the unsaturated cyclic α-angelica lactone, an intermediate formed from the dehydration of LA which results in lower GVL yields. Critically, this work also showed that supported metal catalysts could also afford the further GVL hydrogenation product methyl-THF (MTHF), albeit with modest selectivities of a few percent [12]. Later work by Pinel et al. found that the addition of Re into carbon supported Pd, Pt and Ru catalysts resulted in the further ring opening hydrogenation pathway to yield 1,4-pentanediol [13]. Chia and co-workers further elaborated on the selectivity of RuRe and RuMo supported nanoparticle catalysts through a combination of experimental and computational study. The authors suggested that surface rhenium hydroxyl groups were sufficiently acidic to catalyse the ring opening hydrogenolysis of GVL, which then undergoes further hydrogenation to yield both the mono- and diol products in varying ratios [14].

Given the high activity and selectivity of supported ruthenium catalysts for levulinic acid hydrogenation, considerable work has explored the effect of catalyst preparation conditions on catalyst activity. Palkovits et al. investigated the effect of catalyst support on the activity of impregnation prepared 5 wt% Ru catalysts for LA hydrogenation, showing that the choice of Al₂O₃, SiO₂, TiO₂ or carbon support resulted in a range of LA conversions from 0% to >99%. Interestingly, activity was shown to differ considerably even between commercial support materials obtained from different suppliers. A 5 wt% Ru catalyst supported on a Tronox TiO₂ was shown to be entirely inactive for LA hydrogenation, whilst the Degussa P25-supported analogue exhibited LA conversion and GVL selectivity in excess of 80% under the same conditions [15]. Catalyst support effects were further elucidated by Pan and co-workers, who undertook a similar study of impregnation prepared Ru
catalysts supported on a broader range of metal oxides for the hydrogenation of Levulinic acid, including synthetic zeolites. The authors showed that the metal oxide-supported Ru catalysts were highly selective towards GVL, whilst the use of highly acidic zeolites yielded valerate esters as a result of concerted hydrogenation-esterification with the ethanol solvent [16].

A wide range of catalyst preparation techniques have been used to prepare Ru catalysts active for LA hydrogenation. Impregnation is the most commonly used technique, wherein solution phase metal precursors are adsorbed onto a support material which yields supported metal nanoparticles upon heat treatment. Preparation parameters, such as metal loadings, support materials choice and metal precursor choice have been shown by several groups to produce highly active catalysts. Regalbuto et al. compared the activity of carbon- and alumina-supported Ru catalysts prepared by conventional and selective electrostatic absorption (SEA) impregnation [17]. The SEA-prepared catalysts were shown to be considerably more active than the conventionally prepared analogues, and this was attributed to increased Ru dispersion and a decrease average nanoparticle diameter. The conventionally prepared impregnation catalysts were found to exhibit both a larger average particle diameter and a broader particle size distribution and therefore contained a variety of Ru nanoparticulate species with varying activity.

Colloidal catalyst preparation methods are known to produce highly dispersed metal nanoparticles with narrow particle size distributions [18,19]. Such an approach was used by Jones and co-workers in preparing Ru/C catalysts for LA hydrogenation. Aqueous colloidal Ru nanoparticles stabilized by polyvinyl alcohol (PVA) immobilized on activated carbon were shown to be highly active for LA hydrogenation to GVL, at comparatively mild conditions compared to previous work [20]. Whilst colloidal methods allow for good control of metal nanoparticle size/composition through the addition of stabilizers and chelators, preparations typically take place in highly dilute aqueous solutions to maximize colloidal stability, which generates large amounts of aqueous waste. Furthermore, polymeric stabilizers required during preparation to inhibit nanoparticle agglomeration have been shown to remain on the catalyst surface after preparation and limit reactant access to the active metal nanosurface [21].

The solvothermal decomposition of metal precursors is an alternative method that has shown great versatility in preparing supported metal nanoparticle catalysts. The ‘polyol’ method, so called due to the prolific use of di- and tri-ol solvents, has been used to prepare a plethora of both colloidal and supported metal nanoparticulate materials. The polyol method has been demonstrated to be highly versatile, having been previously used to prepare monometallic Ag, Au, Pd, Pt, Fe, Ni, Co and Sn nanoparticles just to name a few [22–24]. The universal applicability of this method is derived from the mechanism of nanoparticle formation. At elevated temperatures, the solvent forms a highly reductive species, which facilitates the decomposition and reduction of the metal precursors to yield nanoparticles. The reduction potential of these intermediates is sufficiently high to reduce a wide range of transition metal precursors, and the scope of this method can be expanded considerably through the addition of stronger reductants to yield not only metallic nanoparticles but also metal oxides and chalcogenides [25]. Furthermore, the chelating nature of the polyol solvent ensures good solubility of a wide range of precursor compounds and additionally, sterically stabilizes the resultant nanoparticles, minimizing agglomeration.

Typically, polyol syntheses take place at temperatures above 150 °C to afford rapid nanoparticle formation, with preparation temperature depending on the choice of solvent. Traditionally, the required high temperature conditions have been achieved using conduction heating, realized on the laboratory scale using a hotplate. Conduction heating is inefficient and slow, leading to long temperature ramping to reach target temperatures whilst being susceptible to the formation of temperature gradients across reactant mixtures. Microwave heating is an alternative that has emerged over the past two decades in the field of nanomaterials and chemical transformations that offers many benefits over conventional thermal treatments, principally very high temperature ramping rates and minimal thermal gradients [26–28].
Furthermore, microwave heating has been shown to decrease nanoparticle preparation time from hours, in the case of conventional heating, to minutes, and additionally, enables access to highly faceted and crystalline nanomaterials [29]. We have previously shown that AuPd/TiO₂ catalysts could be readily prepared using a microwave assisted solvothermal method in as little as 15 min, allowing for rapid screening of bimetallic nanoparticle catalysts. In this study, we explored the effect of microwave treatment time, temperature and metal precursor on the activity of Ru/TiO₂ catalysts for the hydrogenation of Levulinic acid.

2. Results & Discussion

Catalytic Activity of MW Prepared Ru/TiO₂

Initially, a series of Ru/TiO₂ catalysts was prepared from Ru(acac)₃ at 150, 175 and 200 °C to assess the effect of preparation temperature and time on catalyst activity for LA hydrogenation. The catalysts were denoted in the manuscript using the following nomenclature: reductant-preparation temperature (°C)-preparation time, e.g., Acac-200-10 was prepared using Ru(acac)₃, at 200 °C for 10 min.

The LA hydrogenation activity of the catalysts is presented in Figure 2. For all the catalysts, the sole product observed was γ-valerolactone. It was found that for a given processing time, increasing preparation temperature resulted in an increase in catalyst activity. In the case of the catalysts prepared with 5 min processing time, LA conversion increased from 41% in the case of Acac-150-5 to 51% and 61% for Acac-175-5 and Acac-200-5, respectively. A similar trend was also observed when the preparation time was increased to 10 min, with these catalysts showing higher activity than the 5-min prepared analogues.

![Figure 2](image)

**Figure 2.** Levulinic acid hydrogenation activity of 1 wt% Ru/TiO₂ catalysts prepared using Ru(acac)₃ at different process times and temperatures. Reaction conditions: catalyst (25 mg), 5 wt% LA (10 mL), 100 °C, H₂(5 bar), 1 h.

The longest processing time, 15 min, was found to yield catalysts with relatively poor activity; increasing preparation time from 10 to 15 min resulted in a decrease in LA conversion from 69.4% to only 27% for Acac-200 series. The activities of the catalysts are comparable to previously published 1 wt%Ru/TiO₂ prepared by an aqueous colloidal method [20]. Given the similar catalyst activities, the principle benefit of a microwave assisted solvothermal method is the rapid preparation of highly active catalysts using a minimal amount of solvent, and therefore, highly concentrated metal precursors.
For comparison, preparation of the previously reported sol immobilization 1 wt%Ru/TiO$_2$ catalysts required over 30 times the solvent volume of the analogous microwave prepared catalyst on a per gram basis, principally to avoid colloidal agglomeration.

The weight loading of the catalysts was measured using MP-AES. Due to the insolubility of Ru species in aqua regia, the most commonly used laboratory acid digestion matrix for AES, the weight loadings were instead determined by comparison between the initial concentration of the dissolved Ru precursors pre-preparation and comparing this to Ru concentration measured in the supernatant solvent post catalyst preparation. The results in Table 1 show that the 1 wt% denotation of metal loading was very close to the measured weight loading and that almost all of the metal in the precursor solution was deposited on to the support.

**Table 1.** Theoretical and actual ruthenium loading of Ru/TiO$_2$ catalysts measured by MP-AES analysis of post synthesis supernatent solution.

<table>
<thead>
<tr>
<th>Theoretical Ru Loading (wt%)</th>
<th>Actual Ru Loading (wt%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acac-200-10</td>
<td>1.08</td>
</tr>
<tr>
<td>Acac-200-15</td>
<td>0.97</td>
</tr>
<tr>
<td>Cl-150-5</td>
<td>0.96</td>
</tr>
<tr>
<td>Cl-150-10</td>
<td>1.02</td>
</tr>
</tbody>
</table>

To better understand the relationship between catalyst preparation parameters and activity, the most and least active Ru(acac)$_3$-derived catalysts were investigated by TEM, displayed in Figure 3. The catalysts Acac-200-10 and Acac-200-15 were found to contain highly dispersed Ru nanoparticles, with the former exhibiting a mean particle diameter of 2.35 nm versus 2.55 nm for the latter. The increase in mean particle size could be a result of the sintering of formed nanoparticles or alternatively, surface-induced growth from increased precursor nucleation on the formed nanoparticles. Previous work by Regalbuto et al. investigated the effect of particle size on the activity of alumina and carbon supported Ru nanoparticle catalysts prepared by selective electrostatic adsorption (SEA) for Levulinic acid hydrogenation [17]. The authors showed that the activity of the catalysts correlated with Ru nanoparticle size, yielding a volcano trend with a maximum around 1.5 nm average particle diameter. This contrasts to the structure-activity relationship observed in this work, wherein a modest increase in average particle size of only 0.2 nm between the acac-200-10 and acac-200-15 catalysts was accompanied by a 61% decrease in LA conversion, suggesting that the broad range of catalyst activities presented cannot be ascribed to particle size effects alone.
The two catalysts, Acac-200-10 and Acac-200-15, were further studied by x-ray photoelectron spectroscopy (XPS) to determine whether their respective catalytic activities were a consequence of Ru surface oxidation state variations. The Ru3d/C1s and O1s narrow scan spectra are presented in Figure 4. In the case of the catalyst Acac-200-10, fitting of the Ru3d/C1s spectrum suggests the presence of both a metallic and oxide Ru species with Ru3d5/2 binding energies of 279.5 and 280.4 eV respectively [30]. These binding energies are similar to those reported by Okal and co-workers for analogous Ru/Al2O3 catalysts, with the presence of oxidic Ru species being attributed to surface oxidation between preparation and analysis [31]. The Ru 3d/C1s region also shows that both catalysts contained significant amounts of carbonaceous material, likely a consequence of the preparation method. The XPS derived atomic surface composition, presented in Table 1, shows that both the most and least active Ru(acac)3-derived catalysts exhibit similar surface carbon concentrations, 48.22 and 42.06 at% respectively, suggesting that the formation of carbonaceous deposits is not detrimental to catalyst activity. Research by Skrabalak and co-workers found that in the case of polyol prepared Ag
nanoparticles, both the precursor salts and resultant nanoparticles were highly active for not only the oxidation of the ethylene glycol solvent, but also the oxidation by-products of nanoparticle formation, such as glycolaldehyde [32]. Similarly, Feldmann et al. reported that colloidal $\text{Zn}_3(\text{PO}_4)_2$ nanoparticles prepared by the polyol method exhibited unusual fluorescence activity, which was determined to be due to the dehydration and carbonization of the solvent yielding 3–5-nm carbon dots in addition to the desired metal phosphate nanoparticles [33]. It can therefore be suggested that the high carbon content of the catalysts prepared in this work is a result of either complete carbonization of the solvent and resultant oxidation products during catalyst preparation, or alternatively, due to the formation of large oligomeric condensation products as a result of the condensation reaction also between the solvent and associated oxidation products.

The presence of complex carbonaceous species on the surface of the catalyst is consistent with the C1s XPS contributions at 284.7, 286.2 and 288.6 eV, corresponding to aliphatic C, C-O and C = O environments, respectively [34–36]. Whilst the aliphatic C and C-O components could be indicative of latent ethylene glycol from the preparation procedure, the presence of C = O contributions and high
overall carbon surface concentration is indicative of the formation of carbonaceous deposits resulting from the oxidation of the solvent during catalyst preparation. Similarly, the relative contributions of the three carbon environments and their overall surface concentrations are inconsistent with the previously reported spectrum of solid Ru(acac)$_3$, supporting the hypothesis that the high surface carbon content is due to the formation of insoluble carbonaceous deposits, rather than an incomplete decomposition of the Ru precursor [30]. Fitting of the O1s XP spectrum similarly indicates the presence of three oxygen environments at 529.7, 531.8 and 533.32 eV, consistent with Ti-O, C-O and C=O species, providing further evidence for the formation of oxygcnated carbonaceous deposits.

Given that the XP spectrum of the Acac-200-10 catalyst shows high bulk carbon concentrations, SEM-EDX was undertaken to determine the dispersion of the carbon layer on the catalytic material, as shown in Figure 5. Consistent with the TEM presented in Figure 3, low resolution SEM-EDX mapping of a representative catalyst particle shows that the Ru was well dispersed across the support surface, with a complete absence of larger nanoparticles agglomerates. The EDX C Lα map agreed with the XPS spectra, showing an even distribution of carbon across the surface of the catalyst particle.

![Figure 5. FEG-SEM EDX Map of representative acac-200-10 particle (a) Ru Lα, (b) Ti Lα, (c) FEG-SEM image, (d) O Lα, (e) C Lα. Scale bar representative of 2 μm.](image)

A series of further catalysts were prepared using RuCl$_3$ with a range of preparation temperatures and processing times, given the large variation in activity of the Ru(acac)$_3$-derived Ru/TiO$_2$ catalysts. The LA hydrogenation activity of the catalysts are presented in Figure 6. Unlike the catalysts prepared using Ru(acac)$_3$, the most active RuCl$_3$ catalysts were formed at low temperatures and short processing times. Indeed, the catalyst CI-150-5, prepared at the mildest temperature for the shortest time was the most active catalyst of the series with an LA conversion of 67%. Increasing the microwave irradiation time at 150 °C to 10 or 15 min yielded less active catalysts with LA conversions of 28% and 37% respectively. Likewise, the catalysts prepared at either 175 or 200 °C showed much reduced activity compared to those prepared at 150 °C, with catalyst activity also decreasing with increasing microwave processing time.
The catalysts Cl-150-5 and Cl-150-10 were further studied by TEM, shown in Figure 7. Similar to the Ru(acac)$_3$ derived catalysts, increasing the processing time resulted in an increase in average particle size from 1.87 to 2.26 nm, suggesting that lower temperatures and short processing times may be employed favorably to yield small well-dispersed supported nanoparticles. In the case of the Cl-150-10 catalyst, the particle size distribution was found to be skewed towards smaller particle diameters (<2 nm), whilst larger (>5 nm) particles, which were not found in other samples, were readily observed for this catalyst. This suggests that the decreased activity of the catalysts prepared with long processing times can be attributed to some extent to sintering of the active Ru nanoparticles.
Figure 7. Representative TEM and associated particle size distribution of (a,b) Cl-150-5 and (c,d) Cl-150-10.

XPS Ru 3d/C1s and O1s narrow scans of the RuCl$_3$ derived catalysts Cl-150-5 and Cl-150-10 are presented in Figure 8 and surface elemental compositions in Table 2. Consistent with the Ru(acac)$_3$ derived analogues, the catalysts Cl-150-5 and Cl-150-10 also exhibit very high surface carbon concentrations, suggesting that the formation of insoluble carbon species is independent of the choice of Ru precursor. In addition, peak fitting of the Ru 3d region is consistent with the presence of both Ru(0) and Ru(IV), likely due to the formation of a RuO$_2$ shell in the case of both catalysts.
Figure 8. XPS narrow scans of (a) Cl-150-5 Ru3d/C1s region, (b) Cl-150-5 O1s region, (c) Cl-150-10 Ru3d/C1s region and (d) Cl-150-10 O1s region. Ru3d/C1s region—carbon fitting contributions in greyscale, Ru contributions in red/orange.

Table 2. XPS Derived atomic surface concentrations of 1 wt% Ru/TiO$_2$ catalysts prepared using Ru(acac)$_3$ and RuCl$_3$ at different process times and temperatures.

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>XPS Derived Surface Concentration (at%)</th>
<th>O</th>
<th>Ti</th>
<th>C</th>
<th>Ru</th>
<th>Cl</th>
<th>Ru(0)/Ru(II)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acac-200-10</td>
<td>38.02</td>
<td>13.43</td>
<td>48.22</td>
<td>0.28</td>
<td>0.04</td>
<td>1.3</td>
<td></td>
</tr>
<tr>
<td>Acac-200-15</td>
<td>41.13</td>
<td>16.34</td>
<td>42.06</td>
<td>0.37</td>
<td>0.11</td>
<td>1.4</td>
<td></td>
</tr>
<tr>
<td>Cl-150-5</td>
<td>44.17</td>
<td>18.24</td>
<td>37.16</td>
<td>0.36</td>
<td>0.09</td>
<td>2.6</td>
<td></td>
</tr>
<tr>
<td>Cl-150-10</td>
<td>41.31</td>
<td>16.12</td>
<td>42.18</td>
<td>0.3</td>
<td>0.09</td>
<td>1.5</td>
<td></td>
</tr>
</tbody>
</table>

Given that both the RuCl$_3$- and Ru(acac)$_3$-derived catalysts were found to contain significant surface carbon concentrations by both XPS and SEM-EDX analysis, a series of catalysts was further investigated by thermogravimetric analysis (TGA) under oxidising conditions in an attempt to quantify the amount of carbon present. The results are shown in Figure 9. The mass loss from the catalyst
was found to vary between 2.4% and 4.5% for the Acac-200 and Cl-150 catalyst series. No clear correlation is observed linking catalyst preparation conditions/carbon concentration and catalytic activity. This finding is consistent with the XPS derived atomic surface concentrations presented in Table 2, which shows the presence of high carbon surface concentrations in every catalyst analysed.

The surface area of the catalysts was measured to determine whether the contribution of the carbon overlayer had an effect on the surface area of the catalyst. The results (Table 3) clearly show that no increase in total surface area was observed, and the total surface area of the catalyst remained close to the measured area of the P25 support. Coupled with the SEM results, this suggests that the carbon overlayer formed on the surface of the support material was relatively thin. BET surface area measurements (Table 3) indicate no increase in measures surface area of these catalysts, again indicating that the amount of carbon present is minor compared to the bulk.

### Table 3. BET Surface Areas of Ru Catalysts and P25 Support

<table>
<thead>
<tr>
<th>Material</th>
<th>BET Surface Area (m²/g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>P25 TiO₂</td>
<td>51.6</td>
</tr>
<tr>
<td>Acac-200-10</td>
<td>52.1</td>
</tr>
<tr>
<td>Acac-200-15</td>
<td>52.1</td>
</tr>
<tr>
<td>Cl-150-5</td>
<td>51.8</td>
</tr>
<tr>
<td>Cl-150-10</td>
<td>51.9</td>
</tr>
</tbody>
</table>

3. Materials and Methods

3.1. Catalyst Preparation

An amount of 1 wt% Ru/TiO₂ (1wt% of the total catalyst mass is Ru, 99 wt% TiO₂) catalysts was prepared in a CEM Discover SP microwave reactor using a 35 mL quartz reaction vessel equipped with a magnetic stirrer and sealed with a Teflon lined silicone cap. Stock solutions of Ru(acac)₃ (STREM, Cambs UK) and RuCl₃.xH₂O were prepared in ethylene glycol, yielding solution concentrations of 10 mg Ru/mL. A typical preparation procedure is outlined below.
Briefly, 0.99 g TiO$_2$ (P25, Degussa, Brussels) and 10 mL ethylene glycol (ACROS Organics, UK) were added to the reaction vessel and homogenized with magnetic stirring. An amount of 10 mL Ru stock solution was then added with further stirring. The vessel was then capped and subjected to microwave treatment at 150, 170 or 200 °C for 5, 10 or 15 min. The microwave reactor was operated isothermally, with microwave power output varying dynamically between 0-200 W. After the allotted time, the reaction vessel was cooled and the catalyst slurry filtered and washed using 200 mL distilled H$_2$O. The catalyst was then air dried at 110 °C for 16 h before use.

The catalysts were denoted in the manuscript using the following nomenclature: reductant-preparation temperature (°C)-preparation time, e.g., Acac-200-10 was prepared using Ru(acac)$_3$, at 200 °C for 10 min.

3.2. Levulinic Acid Hydrogenation

The Levulinic acid hydrogenation activity of the catalysts was evaluated in a 50 mL stainless steel autoclave (Parr) equipped with a PTFE liner. The liner was charged with 0.5 g Levulinic acid (Alfa Aesar, Lancs UK), 10 mL H$_2$O (HPLC grade, Fisher Sci.) and 50 mg catalyst. The reactor was then pressurized with 10 barg H$_2$ and heated to the desired temperature, with the reaction initiating with the stirring of the reactor at 1200 rpm. After 1 h, the stirring was ceased, and the reactor cooled with an ice bath. The reactor head space was depressurized with the reactor temperature below 30 °C before the liquid phase was filtered.

The resultant solutions were analyzed using a GC (Varian, Crawley, UK) equipped with a CP-Sil 5CB column (Agilent, UK) and FID detector. The composition of the reaction solutions was determined by calibration against Levulinic Acid and γ-Valerolactone (Alfa Aesar, Lancs UK) standard solutions, in both cases $r^2 > 0.998$.

3.3. Characterization

Scanning electron microscopy was performed on a Tescan (Cambridge, UK) Maia3 field emission gun scanning electron microscope (FEG-SEM) fitted with an Oxford Instruments XMAXN 80 energy dispersive X-ray detector (EDX). Images were acquired using the secondary electron and backscattered electron detectors. Samples were dispersed as a powder onto 300 mesh copper grids coated with holey carbon film.

Transmission electron microscopy (TEM) and scanning transmission electron microscopy (STEM) were performed on a JEOL (Welwyn Garden City, UK) JEM-2100 operating at 200 kV.

X-ray photoelectron spectroscopy (XPS) data was collected on a Kratos (Manchester, UK) Axis Ultra DLD X-ray photoelectron spectrometer using a monochromatic Al Ka X-ray source operating at 140 W (10 mA x 12 kV). Survey scans and high-resolution scans were acquired at a pass energy of 160eV and 20eV respectively. Charge neutralization was achieved using an electron source and spectra subsequently calibrated to the lowest energy C(1s) peak of the adventitious carbon, assigned a binding energy of 284.8 eV; experimental binding energies are quoted ±0.2 eV. Data was processed using CasaXPS (v2.3.23rev 1.0G) using a Shirley background and modified Wagner sensitivity factors as supplied by the manufacturer.

Catalyst metal loading was determined by microwave plasma atomic emission spectroscopy (MP-AES) using an Agilent (UK) MP4100 MP-AES. Briefly, a small aliquot (100 µL) of the pre- and post-preparation solutions was sampled and diluted in 10 wt% aqueous HCl (9.9 mL) prior to analysis. Quantification of the Ru concentration was determined by emission intensity at 372.8 and 349.9 nm, with the emission intensity calibrated using a certified Ru reference standard solution (Agilent).

Brunauer–Emmett–Teller (BET) surface area measurements were performed on a Quantachrome (Hook, UK) Nova 2200 using a multipoint N$_2$ adsorption method. Prior to the analysis, samples were degassed for 24 h at 200 °C under vacuum.

Thermogravimetric Analysis (TGA) of the prepared catalysts was performed using a Perkin Elmer (Pontyclun, UK) TGA4000. The experiments were performed in a synthetic air atmosphere from 30 to 900 °C with a ramp rate of 5 °C/min.
4. Conclusions

In summary, we investigated the effect of catalyst preparation parameters, such as metal precursor, temperature and processing time, on the microwave-assisted one pot polyol synthesis of TiO$_2$-supported Ru nanoparticles. We found that both the temperature and preparation time are critical to producing catalysts active for Levulinic Acid hydrogenation and that even with irradiation times as short as 5 min, the resulting catalysts were found to contain highly dispersed Ru nanoparticles <3 nm in diameter. In each case, the targeted 1 wt% metal loading was achieved and quantified using MP-AES. Interestingly, catalysts prepared with this method were found to contain carbon surface concentrations in excess of 40%, which can be attributed to the formation of insoluble species from the solvent during nanoparticle synthesis. Bulk characterization (SEM-EDX, BET) of the catalysts suggested that the surface carbon layer was comparatively thin and likely spread across the entire catalyst surface. Processing of the catalyst post synthesis to remove this carbonaceous layer, further and exposing the surface of the Ru nanoparticles, could result in even greater activity.


Funding: The authors would like to thank the UK Engineering and Physical Sciences Research Council EPSRC (EP/K031635/1) for financial support. XPS data collection and analysis was performed at the EPSRC National Facility for XPS (‘HarwellXPS’), operated by Cardiff University and UCL, under contract No. PR16195.

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

Data Statement: Information on the data underpinning the results presented here, including how to access them, can be found in the Cardiff University data catalogue at http://doi.org/10.17035/d.2019.0082332629.

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