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Support Induced Control of Surface Composition in Cu–Ni/TiO₂ Catalysts Enables High Yield Co-Conversion of HMF and Furfural to Methylated Furans

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ABSTRACT: 5-(Hydroxymethyl)furfural (HMF) and furfural (FF) have been identified as valuable biomass-derived fuel precursors suitable for catalytic hydrodeoxygenation (HDO) to produce high octane fuel additives such dimethyl furan (DMF) and methyl furan (MF), respectively. In order to realize economically viable production of DMF and MF from biomass, catalytic processes with high yields, low catalyst costs, and process simplicity are needed. Here, we demonstrate simultaneous coprocessing of HMF and FF over Cu–Ni/TiO₂ catalysts, achieving 87.5% yield of DMF from HMF and 88.5% yield of MF from FF in a one pot reaction. The Cu–Ni/TiO₂ catalyst exhibited improved stability and regeneration compared to Cu/TiO₂ and Cu/Al₂O₃ catalysts for FF HDO, with a ∼7% loss in FF conversion over four sequential recycles, compared to a ∼50% loss in FF conversion for Cu/Al₂O₃ and a ∼30% loss in conversion for Cu/TiO₂. Characterization of the Cu–Ni/TiO₂ catalyst by X-ray photoelectron spectroscopy, scanning transmission electron microscopy, and H₂−temperature-programmed reduction and comparison to monometallic Cu and Ni on Al₂O₃ and TiO₂ and bimetallic Cu–Ni/Al₂O₃ catalysts suggest that the unique reactivity and stability of Cu–Ni/TiO₂ derives from support-induced metal segregation in which Cu is selectively enriched at the catalyst surface, while Ni is enriched at the TiO₂ interface. These results demonstrate that Cu–Ni/TiO₂ catalysts promise to be a system capable of integrating directly with a combined HMF and FF product stream from biomass processing to realize lower cost production of liquid fuels from biomass.

KEYWORDS: furfural, hydroxymethyl furfural, hydrodeoxygenation, bimetallic catalysts, biofuels, support effects

1. INTRODUCTION

Lignocellulosic biomass is the most abundant and inexpensive renewable resource that can potentially displace petroleum as a carbon neutral alternative for the production of fungible liquid transportation fuels and commodity chemicals.1−4 However, the production of target chemicals and gasoline range fuels from lignocellulosic biomass has been economically challenging due to the need for multiple processing steps and associated high product yields required in each step.5,6 Biomass is rich in both C₆ (glucan) and C₅ (xylan) polymeric sugars that can be converted by acid-catalyzed dehydration into 5-(hydroxymethyl)furfural (HMF) and furfural (FF), respectively, with high yields.7−11 As such, HMF and FF have been identified as valuable fuel precursors suitable for the production of dimethyl furan (DMF) and methyl furan (MF), respectively, through selective hydrodeoxygenation (HDO) (hydrogenation followed by hydrogenolysis). Due to their high octane numbers, these methylated furans (DMF and MF) have been deemed as valuable target fuel products from biomass suitable for blending with gasoline.12 DMF and MF can also be selectively converted into C₆ and C₅ alcohols as direct gasoline replacements or further converted into long chain hydrocarbons for diesel or jet fuel applications by condensation followed by deoxygenation.13,14 However, catalytic production of DMF and MF from biomass derived HMF and FF requires high yields and process simplicity.

Supported noble metal catalysts have been thoroughly studied for HDO of HMF and FF. To reduce costs and control toxicity from catalyst wastes, heterogeneous base metal catalysts, particularly first row transition metals such as Ni, Cu,
undesired reactions such as decarboxylation and ring hydrogenation are common when using base metal catalysts (particularly Ni, Co, and Fe), and the stability of these materials under reaction conditions is low at high temperatures (>200 °C).18–20 FF conversion to MF occurs through hydrogenation to furfuryl alcohol (FOL) as an intermediate product, followed by hydrogenolysis to MF.21,22 Unwanted side reactions include ring hydrogenation of MF or FOL to form methyl tetrahydrofuran (MTHF), or tetrahydrofurfural alcohol (THFOL) and decarbonylation of FF to form furan, see Scheme 1. Cu catalysts minimize decarbonylation and ring hydrogenation due to their full valence d-band and effectively hydrogenate FF to FOL at low temperatures (<200 °C).23–26 However, the low activity of Cu for H2 activation and stronger interaction of Cu with FF, as compared to FOL, make conversion of FOL to MF on pure Cu catalysts slow.27–29 Cu-based bimetallic catalysts have been proposed to overcome the low reactivity of monometallic Cu for hydrogenation reactions. Various formulations have been proposed, such as Cu–Fe, Cu–Pd, Cu–Cr, and Cu–Ni,26,28,30–34 to increase reactivity or selectivity, but further enhancements in reactivity, selectivity toward MF, and catalyst stability are needed for economical implementation.35

HMF conversion to DMF is known to occur through hydrogenation and hydrolysis of the alcohol and carbonyl groups, but undesirable products are formed through decarboxylation and ring hydrogenation reactions, see Scheme 2. Noble metal catalysts such as Ru, Pt, and Pd and their combination with Cu, Fe, and Co in bimetallic catalysts have been reported for HMF conversion to DMF, although the requirement of Pt group metal catalysts is economically challenging for an industrial process.36–41 Limited reports exist on the successful use of solely first row transitions metals or non-noble metal catalyst systems for high yield conversion of HMF to DMF.42 Recently, it has been demonstrated that modifying Cu catalysts with Zn or Co and graphene enabled high selectivity for DMF production.43 However, in these reports, low reactivity43 and stability,42 or expensive catalyst formulations (graphene),43 limit their commercial viability. Bifunctional Ni catalysts combining dispersed Ni species and acidic supports have also been demonstrated for HMF conversion to DMF with high yields, although these reports are limited by catalyst stability44 or require high H2 pressures.45

It is clear that base metal catalysts are most suitable for HDO of FF and HMF to methyl furans due to their low costs, but limited demonstrations of these catalysts are reported to achieve high reactivity, selectivity, and stability. Furthermore, it has recently been demonstrated that high yield coproduction of HMF and FF can be achieved directly from biomass in a single step process using THF as a cosolvent, thereby enabling integrated downstream catalytic strategies to process a single product stream containing both HMF and FF to reduce overall processing costs.14 However, most previous reports on HMF and FF HDO have considered their catalytic conversion separately. It is expected that coupling base metal catalysts capable of simultaneously converting HMF and FF to methylated furans with recently developed biomass pretreatment technologies could realize significant cost savings for an integrated processing strategy that avoids separating biomass sugar streams.

Scheme 1. Reaction Pathway for FF Conversion

**Scheme 2. Reaction Pathway for HDO of HMF to DMF**

**Scheme 1. Reaction Pathway for FF Conversion**

**Scheme 2. Reaction Pathway for HDO of HMF to DMF**

"FF = furfural, FOL = furfuryl alcohol, MF = methyl furan, THFOL = tetrahydrofurfuryl alcohol, and MTHF = methyl tetrahydrofuran."22

"HMF = 5-hydroxymethyl furfural, MFF = methyl furfural, BHMF = bis(hydroxymethyl) furan, MFOL = methyl furfuryl alcohol, and DMF = dimethyl furan."2
USA) was dissolved in 50 mL of deionized (DI) water and added to 5 g of \( \theta \)-Al\(_2\)O\(_3\) (Inframat Advanced Materials, Catalogue no: 26R-0804UPG, Manchester CT 06042, USA) or TiO\(_2\) (P25, NIPPON AEROSIL Co., LTD, Evonik, Degussa GmbH, Batch No. 4161060398) contained in a round-bottom flask to obtain a 10 wt % loading of Cu. The solution was mixed and dried at 80 °C in a rotary evaporator. Supported Ni catalysts were prepared similarly to Cu catalysts, where nickel(II) nitrate hexahydrate (Ni(NO\(_3\))\(_2\)-6H\(_2\)O, Aldrich, purity 99.99%, Louis, MO 63103, USA) was used as a precursor in desired quantities to achieve 10 wt % loadings on TiO\(_2\) and \( \theta \)-Al\(_2\)O\(_3\). The resulting solids were dried at 100 °C for 12 h in an oven and calcined at 450 °C for 5 h. Prior to reactivity experiments, catalysts were reduced by a pure H\(_2\) flow rate of 50 mL min\(^{-1}\) at 450 °C for 3 h and cooled to 25 °C under the same environment.

### 2.2. Synthesis of Bimetallic Cu–Ni Catalysts

Required amounts of Ni and Cu precursors to achieve 5 wt % loadings of each metal were mixed simultaneously in 50 mL of DI water and added to 5 g of TiO\(_2\) or \( \theta \)-Al\(_2\)O\(_3\) in a round-bottom flask. These materials were then mixed and dried at 80 °C in a rotary evaporator. The solid was collected and dried at 100 °C for 12 h in an oven and calcined by heating at 450 °C for 5 h. Prior to reactivity experiments, catalysts were reduced by pure H\(_2\) at a flow rate of 50 mL min\(^{-1}\) at 450 °C for 3 h and cooled to 25 °C under the same environment.

### 2.3. Catalyst Characterization Techniques

#### 2.3.1. X-ray Diffraction (XRD)

XRD spectra of reduced catalysts were recorded in the 2\( \theta \) range of 20 to 90° on a Micromeritics AutoChem 2920 instrument. In each experiment, 0.1 g of catalyst was placed in a quartz tube and treated with pure Ar flowing at 30 mL min\(^{-1}\) at 150 °C for 1 h. A gas mixture of H\(_2\) (10%)–Ar (90%) was passed through the quartz reactor at 25 °C for 1 h with a 50 mL min\(^{-1}\) flow rate. The temperature was raised to 800 °C at a linear heating rate of 5 °C min\(^{-1}\). A standard CuO powder was used to calibrate H\(_2\) consumption.

#### 2.3.2. Surface Area

The total accessible surface area \( (S_{BET}) \) of the catalysts was measured by \( \text{N}_2 \) physisorption using a Micromeritics ASAP 2020 instrument.

#### 2.3.3. Scanning Transmission Electron Microscopy (STEM)

STEM imaging was performed at 300 kV accelerating voltage on an FEI Titan Themis 300 instrument fitted with an X-FEG electron source, a three lens condenser system, and an S-Twin objective lens. STEM images were recorded with a Fischione Instruments Inc. M3000 High Angle Annular Dark Field (HAADF) Detector at a probe current of 0.2 nA, frame size of 2048 \times 2048, dwell time of 15 \( \mu \)s/pixel, camera length of 195 mm, and convergence angle of 10 mrad. Elemental X-ray microanalysis and mapping were performed utilizing an FEI Super-X EDS system with four symmetrically positioned SDD detectors of 30 mm\(^2\) each, resulting in an effective collection angle of 0.7 sr. Elemental maps were collected in STEM mode with a beam current of 0.4 to 0.25 nA with a 512 \( \times \) 512 pixel frame, dwell time of 30 \( \mu \)s, and acquisition time of up to 10 min. Specimens prepared from suspension in distilled water were deposited on copper grids coated with a lacy carbon. Average metal particle sizes were measured based on the diameter of 100 particles from corresponding TEM images of each catalyst.

#### 2.3.4. X-ray Photoelectron Spectroscopy (XPS)

XPS characterization was carried out using a Kratos AXIS ULTRADLD XPS system equipped with an Al K\( \alpha \) monochromated X-ray source and a 165 mm mean radius electron energy hemispherical analyzer. Vacuum pressure was kept below 3 \( \times \) 10\(^{-9}\) Torr during analysis. Binding energy calibrations were done with reference to the carbon 1s peak by adjusting spectra to 284.8 eV. Depth profiling experiments were conducted by argon sputtering samples for 0, 1, 5, 10, 30, and 60 min with a beam voltage of 4 kV, current of 2.35 A, spot size of 3 \( \times \) 3 mm\(^2\), and vacuum pressure of 3 \( \times \) 10\(^{-9}\) Torr during acquisition. XPS peak fitting for Cu and Ni components was optimized for each support, and parameters of the fit were kept constant. A fwhm of 2 eV (2.15 eV) and a Gaussian/Lorenzian line shape ratio of 30% (60%) was used for all Cu and Ni peak fitting on Al\(_2\)O\(_3\) (TiO\(_2\)). The surface composition of bimetallic Cu/Ni catalysts was calculated using sensitivity factors of 5.321 and 4.044 for Cu and Ni, respectively.

### 2.4. Reactivity Measurements

Prior to each reaction, Ni, Cu, and Cu–Ni catalysts were reduced at 450 °C for 3 h. Without exposure to air, 0.3 g of reduced catalysts was transferred into a 100 mL stainless-steel Parr micro batch reactor containing 1 g of FF (Sigma-Aldrich, 99.9% pure) with 25 mL of either isopropyl alcohol (Fischer Chemical, HPLC grade) or 1,4-dioxane (Fisher Chemicals, HPLC grade) as a solvent. The reactor was initially flushed with H\(_2\) and then pressurized under pure H\(_2\) environments. Next, the reactor temperature was raised to 25−240 °C, and the reaction was conducted for 1−8 h. An identical protocol was used for the HMF and FF/HMF coprocessing reactions, except in HMF coprocessing reactions, 0.5 g of HMF was used as a reactant, and for FF/HMF coprocessing reactions, 0.5 g of FF and 0.25 g of HMF were used.

### 2.5. Product Analysis

Liquid products were analyzed by gas chromatography (Agilent Technologies 7890A; column: DB-WAX Ultra Inert, 30 m long \( \times \) 0.320 mm internal diameter \( \times \) 0.5 \( \mu \)m) via FID according to the following program: hold for 1 min at 30 °C; increase from 30 to 100 °C at a ramp rate of 10 °C min\(^{-1}\); 2 min hold at 100 °C; increase from 100 to 250 °C at a ramp rate of 25 °C/min; 0 min hold; increase from 250 to 325 °C at a ramp rate of 25 °C min\(^{-1}\); and 1 min hold at 325 °C. Mass yields of the final product were quantified by using calibration curves of standard samples in the gas chromatograph. Mass balances accounting for \( >95\% \) of the carbon content were obtained in all experiments. Reactant conversion and product yield were calculated as follows:

\[
\text{HMF (or) FF conversion} \% = \left( 1 - \frac{\text{moles of unreacted FF}}{\text{moles of FF before reaction}} \right) \times 100
\]

\[
\text{Yield} = \frac{\text{moles of the product produced}}{\text{moles of HMF (or) FF before reaction}} \times 100
\]

### 2.6. Catalyst Recyclability

A total of 0.3 g of freshly reduced catalysts was transferred into a 100 mL stainless-steel Parr reactor containing 1 g of FF and 25 mL of 1,4-dioxane. The reactor was pressurized with H\(_2\) to 25 bar, and the reaction was conducted for 2 h at 200 °C. The reactor was cooled by quickly lowering it into a room temperature water bath (25 °C) and then depressurized. The catalyst was separated by filtration, dried at 100 °C for 3 h, and then reused in four recycle experiments without reduction or reactivation. Regeneration
3. RESULTS

3.1. Catalyst Characterization. To explore the influence of support and the formation of bimetallic particles on HDO performance, monometallic Cu (10% weight loading) and Ni (10% weight loading) catalysts and bimetallic Cu−Ni catalysts (5% Cu−5% Ni weight loading) on TiO2 and θ-Al2O3 were synthesized via an impregnation approach. XRD spectra of reduced monometallic (Cu and Ni) and bimetallic (Cu−Ni) catalysts are shown in Figure S1. Reduced monometallic Cu catalysts showed distinct peaks at 43.3° and 50.4° that correspond to the (111) and (200) reflections of metallic Cu, respectively. The reduced monometallic Ni catalysts exhibited clear peaks associated with the (111) reflection of metallic Ni at 44.9°, although in the case of the Ni/Al2O3 catalyst this peak overlapped with Al2O3 reflections. The Cu−Ni/Al2O3 catalyst exhibited a diffraction peak at 43.9°, and the Cu−Ni/TiO2 catalyst exhibited a diffraction peak at 44.1° and a shoulder at 44.6°. The existence of diffraction peaks between the Cu (111) and Ni (111) reflections for the bimetallic catalysts are evidence of the formation of Cu−Ni alloy phases. However, we refrain from using these peak positions to draw conclusions about the nature of the Cu−Ni alloy, as the diffraction peaks are predominantly derived from the largest particles in each catalyst and are not representative of the composition of all particles in each sample.

Table 1. Physicochemical Properties of Monometallic Cu, Ni, and Bimetallic Cu−Ni Catalysts Supported on Al2O3 and TiO2

<table>
<thead>
<tr>
<th>sample</th>
<th>specific surface area (S_{BET}, m²/g)</th>
<th>particle size distribution from TEM (nm)</th>
<th>H₂-uptake (TPR; mmol/g)</th>
<th>Cu⁺/Cu⁰</th>
<th>Ni²⁺/Ni⁰</th>
<th>Cu/Ni (Cu⁰ + Cu²⁺)/(Ni⁰ + Ni²⁺)</th>
</tr>
</thead>
<tbody>
<tr>
<td>θ-Al2O₃</td>
<td>79.7</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>TiO₂ (P25)</td>
<td>54.3</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cu (10 wt %)/Al₂O₃</td>
<td>64.0</td>
<td>7.0 ± 1.6</td>
<td>1.29</td>
<td>37.6/62.4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cu (10 wt %)/TiO₂</td>
<td>43.4</td>
<td>8.0 ± 3.5</td>
<td>1.48</td>
<td>47.8/52.2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ni (10 wt %)/Al₂O₃</td>
<td>64.8</td>
<td>6.4 ± 5.1</td>
<td>1.68</td>
<td>52.4/47.7</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ni (10 wt %)/TiO₂</td>
<td>47.2</td>
<td>9.9 ± 2.6</td>
<td>1.75</td>
<td>57.5/42.7</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cu (5 wt %)−Ni (5 wt %)/Al₂O₃</td>
<td>67.4</td>
<td>4.9 ± 1.6</td>
<td>1.62</td>
<td>38.6/61.4</td>
<td>73.7/26.3</td>
<td>48.7/51.3b</td>
</tr>
<tr>
<td>Cu (5 wt %)−Ni (5 wt %)/TiO₂</td>
<td>45.9</td>
<td>7.5 ± 3.9</td>
<td>1.56</td>
<td>33.1/66.9</td>
<td>57.5/42.5</td>
<td>82.4/17.6b</td>
</tr>
</tbody>
</table>

Table 1. Physicochemical Properties of Monometallic Cu, Ni, and Bimetallic Cu−Ni Catalysts Supported on Al₂O₃ and TiO₂

- Catalysts were calcined prior to analysis. Cu/Ni ratio is from XPS shown in Figure 2, and these are consistent with the values in Figure 3.

was executed via calcination at 450 °C for 5 h followed by reduction with pure H₂ at 450 °C for 3 h.

3. RESULTS

3.1. Catalyst Characterization. To explore the influence of support and the formation of bimetallic particles on HDO performance, monometallic Cu (10% weight loading) and Ni (10% weight loading) catalysts and bimetallic Cu−Ni catalysts (5% Cu−5% Ni weight loading) on TiO₂ and θ-Al₂O₃ were synthesized via an impregnation approach. XRD spectra of reduced monometallic (Cu and Ni) and bimetallic (Cu−Ni) catalysts are shown in Figure S1. Reduced monometallic Cu catalysts showed distinct peaks at 43.3° and 50.4° that correspond to the (111) and (200) reflections of metallic Cu, respectively. The reduced monometallic Ni catalysts exhibited clear peaks associated with the (111) reflection of metallic Ni at 44.9°, although in the case of the Ni/Al₂O₃ catalyst this peak overlapped with Al₂O₃ reflections. The Cu−Ni/Al₂O₃ catalyst exhibited a diffraction peak at 43.9°, and the Cu−Ni/TiO₂ catalyst exhibited a diffraction peak at 44.1° and a shoulder at 44.6°. The existence of diffraction peaks between the Cu (111) and Ni (111) reflections for the bimetallic catalysts are evidence of the formation of Cu−Ni alloy phases. However, we refrain from using these peak positions to draw conclusions about the nature of the Cu−Ni alloy, as the diffraction peaks are predominantly derived from the largest particles in each catalyst and are not representative of the composition of all particles in each sample.

Representative TEM images of the monometallic and bimetallic catalysts are shown in Figure 1a,b and Figure S2, with corresponding particle size distributions shown in Figure S3 and average particle sizes reported in Table 1. The average metallic particle diameter in all catalysts was relatively consistent, with values between 4.9 and 9.9 nm. It was observed that for a given metal composition, the Al₂O₃ supported catalysts exhibited a ~1−2 nm smaller average particle diameter compared to the TiO₂ supported catalysts. This is likely due to the ~20 m²/g greater total surface area of
the Al2O3 supported catalysts compared to the TiO2 supported catalysts, see Table 1. From the TEM analysis, we conclude that the active metal particle sizes in all catalysts are relatively consistent, and significant differences in catalyst performance as a function of support for a given metal composition are not expected to be derived from particle size effects. Elemental mapping executed for the Cu–Ni/TiO2 and Cu–Ni/Al2O3 catalysts shown in Figure 2b–d and f–h demonstrated that Cu and Ni signals coexisted for all observed metal particles. This is consistent with the XRD results, providing evidence that bimetallic Cu–Ni particles formed on both Al2O3 and TiO2.

XPS was used to characterize the oxidation state and surface composition of the monometallic and bimetallic catalysts by analyzing the Cu and Ni 2p3/2 peaks. Because these experiments were performed ex situ, the exposure to air when catalysts were transferred from the reduction reactor to the XPS chamber caused partial metal oxidation. Figure 2a–d shows the Cu 2p3/2 spectra for prereduced monometallic Cu and bimetallic Cu–Ni catalysts on Al2O3 and TiO2. All reduced Cu containing catalysts show signatures of Cu0 with binding energies of 932.1–932.4 eV and Cu2+ with binding energies of 933.7–934.0 eV. Comparing the Cu0 2p3/2 peaks in the monometallic Cu catalysts (0.7 eV versus 0.3 eV), with the observed and well-known stronger interactions between Ni and Cu2+ TiO2, with subsequent formation of CuO species and charge transfer from Cu0 to CuO. All supported Ni catalysts showed three peaks at 852.2 and 853.0 eV, 854.5 and 854.8 eV, and 856.2 and 856.6 eV that are assigned to metallic Ni0, NiO, and Ni(OH)2, respectively. Binding energies for the Ni0 2p3/2 peaks were in the order Ni–Cu/Al2O3 > Ni–Al2O3 > Ni–TiO2 > Ni–Cu/TiO2, with the well-known strong interactions between Ni and TiO2 driving charge transfer from Ti4+ to Ni d-states. The shift in binding energy of the Ni0 2p3/2 peak when comparing the monometallic Ni catalysts on Al2O3 and TiO2 was more significant than that observed for the Cu0 2p3/2 peak in the monometallic Cu catalysts (0.7 eV versus 0.3 eV), demonstrating stronger interactions and increased charge transfer at the Ni/TiO2 interface.

The Cu/Ni surface composition ratio was calculated for bimetallic catalysts after reduction and calcination by summing all contributions to the Cu and Ni 2p3/2 spectra and correction for XPS sensitivity factors. For the Al2O3 supported bimetallic catalyst, an almost equal Cu/Ni surface composition ratio of 48.7:51.3 (Table 1) was observed for the reduced catalyst, consistent with the equal weight loadings of Cu and Ni, the miscibility of Cu and Ni, and their expected nonspecific interactions with Al2O3. Interestingly, the surface composition for the TiO2 supported bimetallic catalyst was significantly enriched in Cu, with a Cu/Ni ratio of 82.4:17.6. As shown in Table 1 and Figures S4 and S5, the bimetallic catalyst surface compositions were almost identical after reduction and calcination. The surface enrichment of Cu and weaker Cu–TiO2 interactions observed by XPS in the Cu–Ni/TiO2 catalyst compared to monometallic Cu/TiO2 combined with the observed and well-known stronger interactions between Ni and TiO2 compared to Cu and TiO2 suggest that preferential interactions between Ni and TiO2 drive the formation of core–shell type structures on Cu–Ni/TiO2 catalysts where Cu is preferentially exposed at the surface.

To further substantiate the conclusion that TiO2 induced the formation of core–shell type structures for the bimetallic catalysts, depth profiling XPS experiments were conducted for reduced Cu–Ni/TiO2 and Cu–Ni/Al2O3 catalysts by using Ar ion bombardment, Figures 3 and S6 and S7. For the Cu–Ni/Al2O3 catalyst, an almost constant 50%:50% composition ratio was observed at all Ar sputtering times, which suggests an even distribution of Cu and Ni throughout the bimetallic particles.
For the Cu−Ni/TiO₂ catalyst, the relative Cu composition decreased from 79.1% to 45.5% as the Ar sputtering time was increased from 0 to 60 min. The switch in composition from predominantly Cu to slightly Ni enriched with increasing Ar sputtering time (and thus depth into the particle) is direct evidence of a core−shell like structure for the TiO₂ supported bimetallic catalysts. Because these catalysts are high surface area materials, a precise relationship between Ar sputtering time and sputtering depth to provide quantitative compositional structure of the particles would be difficult. Regardless of this limitation, contrast between the relative Cu and Ni compositions as a function of Ar sputtering time for the Al₂O₃ and TiO₂ catalysts is strong evidence that TiO₂ induced the formation of core−shell structures where the catalytic surface is Cu enriched and the TiO₂ interface is Ni enriched.

To corroborate inferences regarding metal−support interactions and Cu surface enrichment in Cu−Ni/TiO₂, temperature-programmed reduction (TPR) spectra were measured as shown in Figure 4. The amount of H₂ consumption in the TPR experiments was calculated for all samples, showing relatively consistent values between 1.3 and 1.7 mmol/g catalyst, Table 1, which corresponds to ~8−10% of the catalyst mass. This is in agreement with the nominal ~10% metal loading in all samples, suggesting that nominal weight loadings well represent the actual weight loadings and that most Cu and Ni species were reduced in the TPR experiments.

Figure 4a,b show that the TPR spectra of Cu/Al₂O₃ exhibited a single reduction peak at 191 °C, whereas two peaks were observed for Cu/TiO₂ at 127 and 192 °C. The low temperature reduction peak for Cu/TiO₂ is attributed to CuO species directly interacting with the TiO₂ support, whereas the ~190 °C reduction peak is assigned to bulk-like CuO. In the case of Ni/Al₂O₃, Figure 4c shows that two reduction peaks were observed at 350 and 500 °C. The former is assigned to the reduction of amorphous NiO species, while the latter is assigned to the reduction of crystalline NiO. Depending on the synthesis procedure and calcination temperature, non-stoichiometric and stoichiometric Ni-aluminates may also form, although these species (TPR peaks >500 °C) were not observed here. The TPR spectra of Ni/TiO₂ in Figure 4d showed three peaks at 250 °C, 350 °C, and 450 °C. The peaks at 250 and 350 °C are assigned to strongly interacting amorphous and crystalline NiO on TiO₂ due to their significant shift down in temperature compared to Ni/Al₂O₃, and the peak at 450 °C is assigned to the onset of TiO₂ reduction. TPR results from the monometallic catalysts are in agreement with XPS results, providing evidence for significant TiO₂ interactions with Cu and Ni and for this interaction being stronger for Ni-TiO₂.

For Cu−Ni/Al₂O₃, the reduction peaks associated with Cu and Ni in Figure 4e both shifted to lower temperatures, as compared to the monometallic catalysts supported on Al₂O₃. This result suggests an intimate interaction between Ni and Cu species, consistent with the TEM and the XPS analyses that showed similar surface concentrations of Cu and Ni. For Cu−Ni/TiO₂, the sharp reduction peak observed at 180 °C in Figure 4f was assigned to CuO, reduction, and the very weak reduction peak at 277 °C was assigned to amorphous NiO reduction. The dominant signature of CuO reduction coupled with the lack of peak associated with reduction of CuO species that are directly interacting with TiO₂ (seen previously at 127 °C for Cu/TiO₂) are strong evidence of the core−shell structure for Cu−Ni/TiO₂, where Cu is exposed at the surface due to strong Ni−TiO₂ interactions.

To summarize catalyst characterization, monometallic Cu and Ni catalysts on Al₂O₃ and TiO₂ exhibited clear signatures of metal−support charge transfer on TiO₂, and the strength of this interaction was greater for Ni compared to Cu. For the bimetallic catalysts, STEM imaging and elemental mapping showed that all metal particles observed on TiO₂ and Al₂O₃ contained Cu and Ni, demonstrating the formation of bimetallic particles. For Cu−Ni/Al₂O₃, it was observed that the bimetallic particles contained a homogeneous mixture of Cu and Ni throughout the particles. However, for Cu−Ni/TiO₂ strong Ni−TiO₂ interactions induced the formation of core−shell like structures enriched in Cu at the catalytic surface. While it is well documented that surface segregation in bimetallic catalysts can be driven by reactant adsorption, the use of preferential metal−support interactions to control surface compositions in bimetallic catalysts is much less common.

3.2. Catalytic Activity. 3.2.1. Hydrogenation of Furfural. The conversion of FF to MF follows a consecutive reaction pathway with many parallel reactions that can drive formation of undesired products. In addition to catalyst composition, it has been observed that reaction selectivity (and ultimately...
yields) is sensitive to reaction temperature, time, H₂ pressure, and catalyst/reactant loading. We screened reaction conditions of 180 to 240 °C, 15 to 55 bar H₂, and 5 to 30% catalyst/reactant mass loading ratios to identify the conditions at which both Ni and Cu metals produced the highest MF yields, shown in Figure 5 and Tables S1–S3. It was identified that at a

temperature of 200 °C, 25 bar of H₂ pressure, and a catalyst/reactant loading of 30%, decarbonylation and ring opening reactions were significantly suppressed, and the reactions on all catalysts were mostly by hydrogenation and hydrogenolysis of FF. All subsequent reactions were performed under these conditions. It is also worth noting that for initial catalyst screening, isopropyl alcohol was used as a solvent. However, because of significant solvent conversion, all subsequent reactivity comparisons used 1,4-dioxane as the solvent due to its stability under optimized reaction conditions.

The conversion of FF and yield of various products over monometallic Ni and Cu on Al₂O₃ and TiO₂ catalysts at 1 and 8 (4 for Ni) hour reaction times is shown in Figure 6. Generally, monometallic Cu catalysts were selective for MF production, while Ni catalysts were selective for production of THFOL. sixty The Ni catalysts were significantly more reactive than Cu catalysts, as evidenced by the ~1 order of magnitude longer reaction time required for full FF conversion on Cu. On the Ni/Al₂O₃ catalyst, FF was completely converted with a 21.5% yield of FOL and 17.2% MF at 1 h, while for Cu/TiO₂, 25.7% of FF was converted to yield 12.1% FOL and 12.7% MF at 1 h. Further extending reaction times to 8 h resulted in complete FF conversion for both catalysts with similar final MF yields of 74.9% for Cu/Al₂O₃ and 75.9% for Cu/TiO₂. THFOL yields were suppressed on Cu catalysts compared to Ni, due to repulsive interactions between the Cu 3d band and the aromatic furan ring. To more effectively compare FF conversion on the monometallic Cu catalysts, Figure S8 shows FF conversion and product yields over time on both the Cu/Al₂O₃ and Cu/TiO₂ catalysts. In both cases, the time dependent yield profile of FOL strongly suggests that this species is an intermediate in the production of MF, as previously reported.32,65 There were slight differences in the time dependent yields of FOL and MF for the Cu/TiO₂ and Cu/Al₂O₃ catalysts, particularly at all times <6 h for which higher yields of FOL are realized on Cu/Al₂O₃ than on Cu/TiO₂. These results suggest that the initial hydrogenation of FF to FOL occurs more effectively on Cu/Al₂O₃ whereas the hydrogenolysis of FOL to form MF occurs more readily on Cu/TiO₂. Although Cu/Al₂O₃ is more active than Cu/TiO₂, small amounts of undesired products such as diols and pentanols were obtained at an 8 h reaction time on Cu/Al₂O₃. Differences in reactivity of the monometallic Cu catalysts were likely caused by interactions between Cu and the support, which can change the charge state of the active metal or introduce interfacial reaction pathways and the inherent acidity of the support.44,45 Comparing Ni and Cu catalysts, it is clear that Ni promotes ring hydrogenation to form THFOL, while Cu is selective for hydrogenolysis to form the desired product MF and that the support composition only mildly influenced catalyst performance.

While the nature of the support induced relatively small changes in the reactivity of monometallic Cu and Ni catalysts, the support significantly influenced the selectivity and reactivity of FF conversion for the Cu–Ni bimetallic catalysts, as shown.

Figure 5. FF conversion and product yields for Cu/Al₂O₃ catalysts for the reaction conditions shown along the x-axis at four catalyst loadings. All reactions except those noted by a, b, and c were executed at a FF loading of 1 g, 25 mL of IPA solvent, and a Cu(10 wt %)/Al₂O₃ catalyst loading of 0.05 g. The catalyst loadings for the other three were (a) 0.15 g for Cu (10 wt %)/Al₂O₃, (b) 0.3 g for Cu (10 wt %)/Al₂O₃, and (c) 0.3 g for Cu (25 wt %)/Al₂O₃.

Figure 6. FF conversion and product yields over monometallic Ni and Cu supported on Al₂O₃ and TiO₂ catalysts at different reaction times. All reactions were run with an FF loading of 1 g, a catalyst loading of 0.3 g, 25 mL of 1,4-dioxane as a solvent, a temperature of 200 °C, and an H₂ pressure of 25 bar.

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in Figure 7. Cu−Ni/Al₂O₃ was more active than Cu−Ni/TiO₂ with 100% FF conversion achieved in 0.5 h reaction time for Cu−Ni/Al₂O₃, whereas 8 h was required for complete FF conversion on Cu−Ni/TiO₂. On Cu−Ni/TiO₂, MF and FOL were the primary products, with MF selectivity increasing with reaction time to ultimately achieve a 92.1% yield of MF at 8 h. Conversely, the Cu−Ni/Al₂O₃ catalysts favored furan ring hydrogenation, which resulted in increasing THFOL yields with a subsequent loss of FOL, as reaction time was extended.

When comparing results with Cu−Ni/Al₂O₃ to those with monometallic Ni and Cu catalysts, both the reactivity (time for complete FF conversion) and product distribution (primarily THFOL, MF, and at short times FOL) strongly resembled the behavior of the monometallic Ni catalysts. Cu−Ni/TiO₂ showed reactivity behavior that was more similar to that of Cu, with only MF and FOL as significant products. However, Cu−Ni/TiO₂ exhibited about 2-fold higher conversion at the same reaction time compared to monometallic Cu/TiO₂ and significantly enhanced MF yields at longer reaction times (92% vs 75.9%).

Issues previously identified for the application of Cu-based catalysts to FF HDO are the potential for Cu sintering or carbon deposition affecting stability at reaction conditions. To examine these effects, recycle and regeneration experiments were performed where four sequential reactivity experiments were executed without treating the catalyst between experiments, followed by calcination and reduction of the catalysts prior to a final reactivity experiment. It is expected that a loss in reactivity during the four sequential experiments could be due to carbon deposits and Cu sintering, while regeneration should predominantly remove carbonaceous deposits, allowing differentiation of the catalyst degradation mechanisms. As shown in Figure 8a, using the Cu/Al₂O₃ catalyst, FF conversion dropped from 56% to 10% by the fourth reactivity experiment (R4) and then increased to ~47% after a regeneration step (R5), whereas MF selectivity was relatively similar in the first (R1) and fifth (R5) experiments, Figure 8a. The results are consistent with previous studies of Cu/Al₂O₃ catalysts showing low stability under FF HDO conditions. For Cu/TiO₂, the FF conversion decreased from 47% to 18% by R4 but increased to 37% following regeneration (R5), with a similar selectivity to R1, Figure 8b. The lower loss of reactivity during the four sequential experiments (R1−R4) for Cu/TiO₂ compared to Cu/Al₂O₃ suggests that the amount of carbonaceous deposits is reduced by the use of TiO₂ as a support. However, the similar change in reactivity comparison of R1 and R5 for Cu/TiO₂ and Cu/Al₂O₃ suggests that Cu sintering similarly occurs for both catalysts.

For the Cu−Ni/TiO₂ catalyst, FF conversion decreased from 93.5 to 85.1% from R1 to R4, demonstrating enhanced stability compared to the monometallic Cu catalysts, Figure 8c. Regeneration of Cu−Ni/TiO₂ restored the MF and FOL yields seen in the initial experiment, although a small amount of THFOL formation was also observed. The results demonstrate that Cu−Ni/TiO₂ catalysts exhibit significantly increased resistance to reactivity degradation caused by carbon deposition and sintering compared to Cu/TiO₂ and Cu/Al₂O₃. However, it was also observed that a combination of exposure to reaction conditions
conditions and regeneration of the Cu–Ni/TiO$_2$ catalyst resulted in the exposure of small amounts of surface Ni domains that drive ring hydrogenation.

3.2.2. HDO of HMF and Coprocessing of FF and HMF. Because Cu–Ni/TiO$_2$ showed excellent reactivity, MF selectivity, and stability for FF conversion, this catalyst was further tested for reactivity in HMF conversion to DMF and simultaneous coprocessing of HMF and FF to DMF and MF, respectively. The measured time dependent conversion of HMF and HMF with FF (coprocessing) and product yields on Cu–Ni/TiO$_2$ are shown in Figure 9a and b, respectively. HMF conversion to DMF is known to occur through two reaction pathways. In the first pathway, hydrogenolysis of HMF yields methyl furfural (MFF) and then hydrogenation followed by hydrogenolysis of MFF to DMF, as shown in Scheme 2. In a second pathway, 2,5-bis(hydroxymethyl) furan (BHMF) is the intermediate product, and further hydrogenation of both C–O groups in BHMF results in DMF formation. The conversion of either BHMF or MFF to DMF typically occurs through methylfurfural alcohol (MFOL), per Scheme 2. However, MFOL is highly reactive and immediately converts into DMF by hydrogenolysis.

For HMF conversion over Cu–Ni/TiO$_2$, 95.9% conversion of HMF was observed within 0.5 h of reaction, with the primary products being DMF (52.2% yield) and MFF (30.7% yield). With increasing reaction time, DMF was converted selectively to DMF to ultimately result in 84.3% yield of DMF at 8 h of reaction time. Small amounts of MFOL were also observed as a rapidly converted intermediate in the production of DMF from MFF.

In the coprocessing of HMF and FF over Cu–Ni/TiO$_2$, 0.5 g of FF and 0.250 g of HMF were used as reactants with 0.3 g of the catalyst under the same reaction conditions as the HMF and FF cases. Cu–Ni/TiO$_2$ showed similar activity and methylated furan (MF + DMF) yields as in individual processing of HMF and FF. Moreover, based on the time dependent yields of intermediate products (FOL and MFF, for example), the mechanism of methylated furan production did not change for coprocessing of FF and HMF. The maximum yields of MF (88.5%) and DMF (87.5%) were realized after 8 h of reaction at very similar yields to those for the individual processing cases.

Summarizing the reactivity results, the monometallic Cu and Ni catalysts showed FF hydrogenation and HDO reactivity consistent with previous reports, and their reactivity exhibited minimal sensitivity to support composition. However, the bimetallic Cu–Ni catalysts showed significant support effects for FF HDO. On Al$_2$O$_3$, the bimetallic catalyst acted similarly to Ni alone, while on TiO$_2$, the Cu–Ni bimetallic catalyst enhanced rates and MF selectivity compared to Cu alone. The Cu–Ni/TiO$_2$ catalyst showed good stability, regenerability, reactivity, and MF selectivity and outperformed monometallic Cu catalysts in all performance metrics under similar conditions. Finally, Cu–Ni/TiO$_2$ showed excellent selectivity toward methylated furans in HMF and FF/HMF coprocessing reactions.

4. DISCUSSION

The significant support effect observed for bimetallic Cu–Ni catalysts can be explained and rationalized through the catalyst characterization studies. Cu–Ni/Al$_2$O$_3$ reactivity studies suggested that the behavior of the catalyst significantly resembled pure Ni catalysts, where ring hydrogenation was prevalent. Both TPR and XPS showed evidence of equal amounts of Cu and Ni exposed at the catalyst surfaces, while the TEM images suggested that all catalytic particles contained Ni and Cu. On Ni, the primary η$^5$(C==O) FF adsorption geometry brings the aromatic ring close to the surface, enabling ring hydrogenation (Figure 10). Because the ring-surface interaction is only expected at Ni surface domains that have Ni ensemble sizes large enough to coordinate to the furan ring, or at Cu–Ni alloy surface structures with electronic structures significantly resembling monometallic Ni, it can be concluded that Ni-rich domains at the surface of bimetallic Cu–Ni particles were responsible for a majority of the reactivity of the Cu–Ni/Al$_2$O$_3$ catalyst. The proposed Cu–Ni bimetallic particle structure on Al$_2$O$_3$ based on the XPS measurements and reactivity results is shown schematically in Figure 10.

The Cu–Ni/TiO$_2$ catalyst reactivity significantly resembled that of Cu alone, albeit with enhanced reaction rates and MF selectivity. This result can be explained based on the depth profiling XPS, TPR, and TEM results, which together suggest that TiO$_2$ supported bimetallic Cu–Ni particles segregate into core–shell geometries that preferentially expose Cu at their surface, as shown schematically in Figure 10. On Cu surfaces, the FF ring is repelled from the surface due to the full valence d-band, forming the η$^1$(O) FF adsorption geometry, and thus, as opposed to Ni surfaces, carbonyl hydrogenation is favored.
on Cu over ring hydrogenation. The enhanced reactivity and MF selectivity on Cu–Ni/TiO2 compared to monometallic Cu catalysts, with no additional ring hydrogenation or decarbonylation, is primarily attributed to increased rates of H2 dissociation.64,65 The enhanced rates of H2 dissociation but retained Cu selectivity characteristics could occur at so-called single atom surface alloys, where small collections (1–3, essentially small enough so they cannot coordinate to a furan ring) of Ni atoms are exposed at the Cu surface to allow for enhanced H2 dissociation rates but not the η2(C=O) FF adsorption geometry.64,72,73 These reactivity characteristics could also be consistent with a low relative Ni concentration near surface alloy that electronically or sterically modiﬁes the exposed surface Cu atoms and enables enhanced H2 dissociation rates.76,75 Although we cannot differentiate these two mechanisms, it is clear that Cu was significantly enriched at the surface in Cu–Ni/TiO2 catalysts, yet still modiﬁed by Ni in a manner that enhanced H2 dissociation rates without changes in the inherent preference of Cu for carbonyl hydrogenolysis over ring hydrogenation (see Figure 10).

The above discussion, combined with the weak inﬂuence of the support composition on the reactivity of the monometallic catalysts, strongly suggests that the primary role of TiO2 in promoting the catalytic properties of the Cu–Ni catalyst is indirect, with Ni driven to the TiO2 interface and Cu preferentially exposed at the surface. Thus, TiO2 controls the bimetallic Cu–Ni catalyst surface composition but does not directly participate in the catalytic process.76,77 A well-known behavior on TiO2 supported Pt-group metal catalysts is strong metal support interactions (SMSI), where support reduction (oxygen vacancy formation) drives TiO2 encapsulation of the active metal particle.78–80 While a metal selective SMSI encapsulation state that preferentially encapsulates Ni could be imagined, the TPR data oppose this interpretation. SMSI encapsulation states with TiO2 are typically observed following >500 °C reduction treatment. Because minimal Ni reduction peaks were observed for the Cu–Ni/TiO2 catalysts despite Ni reduction peaks being clearly observable in the Ni-TiO2 catalyst, Ni is likely buried subsurface in the bimetallic particles prior to SMSI encapsulation layer formation. This mechanism is further supported by the XPS analysis of surface composition in the prereduced and preoxidized Cu–Ni/TiO2 catalysts that show identical Cu/Ni surface concentration ratios in Table 1.

Thus, preferential interactions between Ni and TiO2 are believed to drive the formation of core–shell like particles where Cu is primarily exposed at the catalyst surface, see proposed structures in Figure 10. We propose that support induced bimetallic particle segregation may be quite general for reducible oxide supported bimetallic catalysts, given known metal speciﬁc interactions with reducible supports.79

The structural properties of Cu–Ni alloys have been extensively studied for their potential application to a range of reactions.81–89 Bulk Cu–Ni alloys are miscible and equilibrated into alloy phases when heated above ~400 °C.90 In bulk systems (polycrystalline ﬁlms, single crystalline ﬁlms, and other structures), Cu surface segregation has been reported to be due to the lower surface energy of Cu in a vacuum compared to Ni.94,91–95 It has further been demonstrated that the surface segregation is a facet dependent phenomenon that occurs more predominately on (100) surface facets compared to (111) surface facets.89,96 Insights into the structure of bulk Cu–Ni alloys have not translated into a complete understanding of the structure of supported Cu–Ni alloy nanoparticles, where multiple surface facets are simultaneously exposed, and the distribution of Cu and Ni is also inﬂuenced by interfacial interactions with the support and environment.

Previous reports of 50%–50% Cu–Ni alloy nanoparticles supported on irreducible supports (SiO2 and Al2O3) have found that as-synthesized particles show no evidence of Cu surface segregation.87,89,97,98 However, reports of Cu–Ni alloy nanoparticles on reducible supports (CeO2, ZrO2, MgO), or supports with signiﬁcant surface defect concentrations (SBA-15), have shown evidence of Cu enriched surfaces, in agreement with our ﬁndings.49,98,99,100 Thus, while bulk Cu–Ni alloys exhibit Cu surface segregation due to the decreased surface energy of Cu in a vacuum, it seems that for supported Cu–Ni alloy nanoparticles, preferential Ni-support interactions on reducible or defective supports is the primary driving force for Cu surface segregation. Also, reconstruction of the as-synthesized catalytic structure under reaction conditions may be expected when signiﬁcant adsorbate–metal speciﬁc interactions exist, for example in a CO atmosphere.90 However, based on the lack of ring hydrogenation observed in our studies for the Cu–Ni/TiO2 catalyst, it can be concluded that migration of Ni to the catalyst surface is minimal under reaction conditions.

It is worth comparing our results to recent reports that examined the inﬂuence of modiﬁcation of Cu catalysts with Ni on reactivity in FF HDO. A surface science approach was used to compare pure Cu (111) and Ni (111) surfaces with a Ni surface layer on Cu (111) and a Cu surface layer on Ni (111) for FF hydrogenation.101 It was clearly seen that in both surface layer conﬁgurations, bimetallics enhanced selectivity toward MF formation compared to monometallics. However, even in the case of Cu monolayers on Ni (111), signiﬁcant nonselective decomposition of FF was observed. This result suggests that for the Cu–Ni/TiO2 catalysts examined here, the Ni concentration in the near surface region is much lower than for the prior surface science study. A separate study of the inﬂuence of Ni addition at various concentrations to Cu/Al2O3 on FF hydrogenation reactivity and selectivity53 showed that increasing the Ni loading increased the rate of FF conversion but also increased the selectivity toward ring hydrogenation, decarboxylation, and ring opening products. The introduction of self-assembled organic monolayers to the catalyst lowered the Ni surface content in Cu–Ni/Al2O3, thereby boosting selectivity.

Figure 10. Schematic diagram of the proposed operating states of Cu–Ni bimetallic catalysts on Al2O3 and TiO2. For Cu–Ni/Al2O3, significant exposure of extended Ni domains drives efﬁcient ring hydrogenation to form THFOL, whereas the segregated structure of Cu–Ni on TiO2 facilitates MF formation. EINT: Energy of interaction between metal (either Ni or Cu) and support (Al2O3 and TiO2).
and reaction rate for hydrogenation products (we note that FOL was the main product in these studies, which is likely caused by the reactions being performed in the gas phase). In this case, it seems that the use of self-assembled monolayers to control the exposed surface concentration of Ni in bimetallic Cu–Ni catalysts bears some resemblance to our reported use of TiO2 to control the Ni surface concentration in bimetallic Cu–Ni catalysts. Although, the Cu–Ni/TiO2 catalysts reported here enable complete conversion and high yields to MF and are expected to have enhanced stability compared to the organic functionalized Cu–Ni catalysts reported previously, thus creating a more scalable approach for controlling Cu surface chemistry for selective hydrogenation reactions. In addition, it was very recently reported that increasing the Ni content in Cu–Ni/MgAl2O catalysts enhanced FF conversion (>99%); however, FOL and THFOL were the only selective products observed.38

In addition to considering how TiO2 induced segregation of Cu–Ni catalyst particles influenced reactivity and selectivity, it is also interesting to explore how this factor may have influenced stability and catalyst regeneration. In the recycling-regeneration experiments shown in Figure 8, it was observed that the Cu/TiO2 and Cu/Al2O3 catalysts exhibited similar performance degradation that is ascribed to carbonaceous deposits on the metal and Cu sintering. The improved stability of Cu–Ni/TiO2 compared to the monometallic catalysts during recycle experiments (R1–R4) is likely due to decreased carbon deposition on the catalysts driven by increased H2 dissociation rates. Increased stability of the Cu–Ni/TiO2 catalyst compared to the monometallic Cu catalysts when considering R1 and R5 is likely due to reduced sintering of the active Cu metal. This suggests that Ni serves as an anchoring site for Cu on TiO2, which enhances catalyst stability and provides a stable platform for regeneration of the core–shell Cu–Ni structure to that for high MF selectivity and reactivity. Further reduction of the Ni loading in Cu–Ni/TiO2 catalysts may allow for similar enhanced reactivity and stability as observed here while also minimizing the surface exposure of Ni observed with increased time under reaction conditions.

Recently, it has been shown that cosolvent-enhanced lignocellulosic fractionation (CELF) of raw biomass enables the production of HMF and FF with extremely high yields.11 Separation of HMF and FF in a liquid stream from the remaining lignin can be effectively achieved through various approaches to yield a combined stream of HMF and FF that is typically rich in FF. Our demonstration of high yield, single pot conversion of FF and HMF over stable and regenerable Cu–Ni/TiO2 catalysts opens new possibilities for an efficient and high yield biomass to the fuel conversion process with only a few required process steps. It is expected that coupling of CELF pretreatment of cellulose biomass with FF/HMF catalytic coprocessing will enable an effective approach for conversion of raw biomass to high quality fuels.

5. CONCLUSIONS

In conclusion, our study showed that Cu–Ni/TiO2 enabled high yield (~90%) conversion of FF and HMF to methylated furans in either single or coprocessing schemes, results not possible with monometallic Cu and Ni, or Cu–Ni/Al2O3. The reactivity of Cu–Ni/TiO2 is proposed to result from strong and selective Ni–TiO2 interactions that favored the formation of the Cu-shell and Ni-core structures, allowing for high selectivity in HDO and enhanced reactivity compared to monometallic Cu catalysts. Furthermore, the strong Ni–TiO2 interactions effectively anchored the bimetallic particles to the TiO2 support, thereby reducing catalyst degradation via sintering and enabling effective regeneration. Finally, it is envisioned that a potentially economical biomass to fuel conversion process can be achieved by coupling CELF pretreatment of raw biomass to produce high yield liquid streams of HMF and FF with catalytic coprocessing using Cu–Ni/TiO2 to high yields of methylated furans.

■ ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acscatal.7b01095.

Additional catalyst characterization (XRD, TEM, XPS) and reactivity data provided in Figures S1–S8 and Tables S1–S3 (PDF)

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