SYNTHESIS OF HMF FROM GLUCOSE IN AQUEOUS MEDIUM USING NIOBIUM AND TITANIUM OXIDES

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ABSTRACT
The effects of H₃PO₄ treatment on the catalytic activity of both niobium and titanium oxides were evaluated during the synthesis of 5-hydroxymethylfurfural (HMF) from glucose in aqueous medium. Catalysts performances were related to acidic and basic properties of oxides. The presence of weak basic sites favored fructose formation via glucose isomerization. The increase in acid strength favored, initially, the formation of 5-hydroxymethylfurfural and, later, its transformation into organic acids and humins. Among the catalysts used in the experiments, phosphated TiO₂ showed the best results, although all catalysts exhibited a significant deactivation along the reaction time.

KEYWORDS
HMF; glucose; oxides; aqueous medium

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1. INTRODUCTION

In past decades, especially in recent years, the interest in petrochemicals and fuels production from biomass has grown substantially. An increase in the societal demand for more sustainable and environmentally responsible products led to the intensification of these activities.

In this context, numerous building blocks are being studied by the scientific community. The 5-hydroxymethylfurfural (HMF) appears among the most promising ones. This compound can be used as a renewable intermediate for the production of different biofuels and monomers in the petrochemical industry as levulinic acid and 2,5 furan dicarboxylic acid (2,5 FDCA) (Lanzafame et al., 2011; Petersen and Werpy, 2004).

Several biofuels like gasoline, diesel, and kerosene, can be synthesized by the 2,5 FDCA esterification or by the HMF etherification with different chain-size alcohols and olefins (Gruter, 2010A; Gruter, 2010B; Gruter and Manzer, 2010A; Gruter and Manzer, 2010B). This process leads to the formation of different furan derivatives with specific and customizable fuel properties such as initial and final boiling point, solidification point, heat value, vapor pressure, along with others.

The HMF can be obtained by hexoses dehydration, especially the fructose (Lewkowski, 2001), as illustrated in Figure 1. Ketohexoses, as fructose, are much more reactive than aldohexoses, as glucose. Thus, ketohexoses get dehydrated by HMF faster than aldohexoses, reducing the formation of side products as organic acids and a tarry oligomeric material called humins. Although the use of glucose as feedstock for HMF production presents some technical difficulties, its low cost, when compared to the use of fructose, makes it utilization competitive (Corma et al., 2007).

Therefore, rising scientific efforts are in place to find novel ways of producing HMF with good yields and in a feasible way. Currently, processes using biphasic systems for continuous extraction of the produced HMF and ionic liquids aiming the reduction of HMF side-products are being developed (Leshkov et al., 2006; Lansalot-Matras and Moreau, 2003; Tong et al., 2010). Meanwhile, financial issues associated with the process of solvent separation and the cost of ionic liquids still represent a challenge.

Figure 1. A typical reaction scheme for HMF production from hexoses.
In this scenario, the aqueous medium displays interesting advantages such as low cost and the inexistence of media toxicity. It also shows high sugar solubility. However, higher HMF yields must be achieved in order to make this process competitive.

The use of homogeneous acid catalysts in aqueous medium was the first attempt studied involving the dehydration of hexoses to produce HMF. Actually, hundreds of different organic and inorganic acids are qualified to dehydrate hexoses (Fan et al., 2011). Nevertheless, problems related to their toxicity, difficulty in separation, and corrosiveness lead to, from a process standpoint, an increased interest broadening in the use of heterogeneous catalysts as zeolites, heteropoly acids, resins, oxides, and others.

Oxides and phosphates are solid acids also used in a large number of reactions such as hydration, esterification, etherifications, dehydrations, and other processes which require a strong acid strength. Niobium acid (hydrated niobium oxide – Nb₂O₅·nH₂O) is known to be a very water tolerant acid catalyst (Okuhara, 2002) and, thus, a very promising catalyst for the dehydration of hexones to HMF. This catalyst was evaluated in the dehydration of fructose in a continuous flow reactor, and the results suggested that HMF decomposition products were not formed under the employed conditions (Cartini et al., 2011).

Niobium phosphate was also evaluated in the dehydration of fructose in water using in a fixed bed reactor (Cartini et al., 2006). In this work, a higher intrinsic effective acidity, in terms of number of acid sites, for the niobium phosphate catalyst was related to a higher initial activity and also to a higher deactivation during the experiments, when compared to the niobium oxide catalyst. Selectivity of around 35% mol/mol was achieved with fructose conversion of 65% mol/mol.

The role of Brönsted and Lewis acid sites in Nb₂O₅ and phosphated Nb₂O₅ was investigated in the transformation of glucose into HMF (Nakajima et al., 2011). In this work, the authors found that Brönsted acid sites have great ability to hydrate formed HMF in water, reducing the HMF yield. In contrast, Lewis acid sites are able to selectively dehydrate glucose to HMF and also to promote the isomerization of glucose into fructose. Indeed, glucose-fructose isomerization may proceed either via hydride transfer (Lewis acid catalyzed reaction) or by proton transfer (basic catalyzed reaction), which is well known in the literature (Souza et al., 2012).

Some metal oxides, for instance TiO₂ and ZrO₂, act as acid and base catalysts for hexoses dehydration in hot-compressed water (Watanabe et al., 2005A). The basic sites are responsible for isomerizing a less reactive feedstock, such as glucose into fructose via proton transfer, while the acid sites catalyze hexoses dehydration into HMF. The ability to isomerize glucose into fructose allows a significant increment in the HMF yield.

TPD measurements were used to evaluate the acidic and basic catalytic properties of TiO₂ (anatase and rutile) and ZrO₂ (monoclinic/tetragonal mixture) on glucose and fructose dehydration in hot compressed water (Watanabe et al., 2005B). Among the three catalysts, TiO₂ anatase showed the highest density of acidic and basic sites while ZrO₂ showed the highest amount of acidic and basic sites. TiO₂ rutile did not show any catalytic activity for glucose dehydration.

In these experiments, TiO₂ anatase showed the highest HMF yield while ZrO₂ showed the highest fructose yield. These results suggest that the density of acid and base sites was probably important for glucose dehydration to HMF, while the amount of weak to moderate basic sites was required for glucose isomerization into fructose.

As mentioned by various researchers (Okazaki et al., 1987; Tang et al.; 2010; Yang et al., 2011), metal oxides treated with H₃PO₄ have a higher density of acid sites and display an enhanced acid strength, which are important in several acid catalyzed reactions.

However, the effect of TiO₂ anatase treated with phosphoric acid on glucose dehydration in hot compressed water aiming the formation of HMF has never been reported in the literature. Thus, the role of the treatment of titanium oxide and niobic acid with phosphoric acid was evaluated on glucose dehydration to HMF in aqueous medium. The catalytic performance was related to the acidic and basic properties through the temperature-programmed CO₂ desorption (CO₂-TPD) and Fourier transform infrared spectroscopy pyridine desorption (pyridine- FTIR).
2. EXPERIMENTAL

2.1 Materials

Glucose (99 wt.%) and HMF (99 wt.%) were purchased from Sigma-Aldrich and Satochem, respectively. Nb₂O₅·nH₂O (99.5 wt.%) was supplied by Companhia Brasileira de Metalurgia e Mineração (CBMM) in powder form (> 99.5 wt.%). TiO₂ anatase (99 wt.%) was purchased from Sigma-Aldrich.

2.2 Catalysts preparation

The phosphated niobium (Nb₂O₅·P) was prepared according to the method described elsewhere (Yang et al., 2011) by mixing 10g of Nb₂O₅·nH₂O in 135mL of 1M H₃PO₄. The suspension was stirred for 52 hours at room temperature. The suspension was aged for 8 hours and, then, filtered. Following, the precipitate was washed several times with deionized water until reaching a neutral pH. After filtration, the powder was dried in vacuum at 60°C for 18 hours and, then, calcinated in air at 400°C for 3 hours.

The phosphated titanium catalyst (TiO₂·P) was prepared using the same procedure. Pure niobic oxide (Nb₂O₅) and anatase titanium oxide (TiO₂) were dried and calcinated under the same conditions.

2.3 Catalysts characterization

Samples surface-areas analyses were measured by nitrogen adsorption at ~196°C using a sorptometer (ASAP-2020, Micromeritics). The samples were degassed at 300°C for 18h prior to measurements. Surface areas were determined by BET method.

X-ray diffraction analyses (XRD) were carried out in a Rigaku Miniflex diffratometer equipped with a copper tube (λ=1.5417Å), operating at 30kV and 15mA and with acquisition intervals of 2θ=0.050, and 2 second per pass.

The basic properties were studied by temperature-programmed desorption of CO₂. CO₂-TPD analyses were carried out in a Balzers Omnistar QMS200 quadrupole mass spectrometer. The samples (~500mg) were pre-treated in situ under He flow (50 mL.min⁻¹) at 400°C for 30 min.

After cooling to room temperature, under He flow, CO₂ was adsorbed for 1 hour at room temperature and with 12.5 mL.min⁻¹ flow. Then, the bulk CO₂ was purged with He for 1 hour at room temperature. The desorption was performed under He flow (50 mL.min⁻¹) until 800°C (10 °C.min⁻¹).

Pyridine-FTIR analyses were performed in a Nicolet Magna 560 spectrophotometer (4000 – 400cm⁻¹ with 4cm⁻¹ resolution). The samples (~30mg) treatments were performed under high vacuum at 400°C for 1 hour, followed by 5 pulses of air at 400°C. Then, the samples were subjected to high vacuum for 30 min. Later, pyridine was adsorbed for 30min at 25°C and 0.266kPa, followed by desorptions under high vacuum at 25°C for 30min, 150°C for 30min, 250°C for 30min and 350°C for 30min.

After pyridine desorption in each temperature, the samples were kept under vacuum and room temperature for 30min before spectrum acquisitions. These spectra were obtained with 4cm⁻¹ resolution after 120 readings at room temperature. Acid-site densities were calculated using 1450cm⁻¹ and 1550cm⁻¹ bands for Lewis and Brønsted acid sites, respectively. Absorptivity coefficients for each band were obtained from available literature (Emeis, 1993).

2.4 Reaction procedure

All the catalytic tests were carried out in a 300mL stainless steel reactor with mechanical agitation and controlled temperature. Prior to reaction, milliQ water was treated in an ultrasound bath for 20 minutes, at room temperature, to remove dissolved oxygen. Then, 2g of d-glucose were dissolved in 198g of degassed water and the catalyst was added in a catalyst/glucose mass ratio of 0.10. Subsequently, a small flow of ultrapure nitrogen was passed through the mixture for 30 minutes to ensure the complete removal of oxygen. Then, the reactor was purged three times with nitrogen, pressurized to 1723.7kPa with the same gas and, later, heated to the reaction temperature. Zero time was considered when temperature reached 200°C. At this time, agitation at 700rpm was turned on. Reaction progress was monitored by taking samples of reaction medium.

2.5 Product analysis

Products and reactants were analyzed by HPLC
In the experiments, glucose conversion ($X_G$), HMF yield ($Y_{HMF}$), selectivity to HMF ($S_{HMF}$), and selectivity to fructose ($Y_F$) are defined as follows:

$$X_G = \frac{nG_0 - nG}{nG_0} \quad (1)$$

$$Y_{HMF} = \frac{nHMF}{nG_0} \quad (2)$$

$$S_{HMF} = \frac{Y_{HMF}}{X_G} \quad (3)$$

$$Y_F = \frac{nF}{nG_0} \quad (4)$$

Where $nG_0$ is the number of moles of glucose initially fed in the reactor; $nG$, $nHMF$ and $nF$ are the number of moles of glucose, and HMF and fructose, respectively, are presented in a given reaction time.

Although levulinic and formic acids could be identified in the chromatograms, their signals and resolutions were very poor, precluding the determination of their yields.

### 3. RESULTS AND DISCUSSIONS

#### 3.1 Surface areas

The catalysts surface areas are shown in Table 1. Surface areas of TiO$_2$ and TiO$_2$-P were similar indicating that treatment with H$_3$PO$_4$ did not modify this property for titanium catalysts. However, Nb$_2$O$_5$-P presented a reduction on its surface area when compared to Nb$_2$O$_5$, phenomenon in accordance with the literature (Okazaki et al. 1987; Tang et al., 2010). The phosphoric acid treatment results in the formation of amorphous niobium phosphate on the Nb$_2$O$_5$ (Tang et al., 2010), which could be responsible for plugging pores and, consequently, reducing surface area.

#### 3.2 X-Ray Diffraction

The catalysts X-ray diffractograms are shown in Figure 2. As one can be observe in Figure 2, the Nb$_2$O$_5$ and Nb$_2$O$_5$-P that calcinated at 300°C were amorphous materials, as reported in the literature (Armaroli et al., 2000; Tang et al., 2010).

Regarding the TiO$_2$ diffractogram, peaks at 25.5° associated to the anatase phase (JCPDS, No. 21-1272) were observed, while no peaks, at 27.45°, were associated with rutile phase for this catalyst.

Concerning the TiO$_2$-P catalyst, no evidence peaks of titanium phosphate, at 27.5°, appeared in the x-ray diffraction (Fan et al., 2008). In
In accordance to the literature, two peaks, at 22.55° and 27.65°, were associated to the crystalline TiP$_2$O$_7$ pattern. These peaks were observed when the phosphated TiO$_2$ is calcinated at 1073°C (JCPDS, No. 38-1468). At lower calcination temperatures, such as 400 and 600°C, there were no records of XRD peaks corresponding to titanium phosphate, suggesting that phosphorus exists as amorphous phosphate (Fan et al., 2008). In the present work, the calcination procedure was carried out at 300°C, thus, no peaks related to phosphated titanium were expected.

### 3.3 CO$_2$ thermodesorption

The CO$_2$-TPD profiles of the calcinated catalysts are shown in Figure 3. Based on the area under each desorption profile, the total desorbed CO$_2$ amount by mass unity and density of basic sites of each catalyst were calculated, and the results are shown in Table 2.

Comparing the niobium catalysts, the Nb$_2$O$_5$ showed a narrower peak at 642°C. The phosphoric acid treatment reduced the basic strength of sites, indicating a broader distribution of these sites. According to the literature, the H$_3$PO$_4$ treatment in metal oxides reduces the electron donation capacity, implying a reduction of basic strength (Samantaray and Parida, 2001). However, the titanium catalysts showed an opposite behavior. The TiO$_2$-P showed a higher CO$_2$ desorption temperature, suggesting that this catalyst has stronger basic sites when compared to the untreated titanium oxide.

The metal oxides basicity increase is usually related to electrons trapped in defects sites, which enhance the catalyst electron donation capacity (Samantaray and Parida, 2001). Although this phenomenon is usually observed at higher temperatures, it could be an explanation for the phosphated titanium oxide anomalous behavior.

### 3.4 FTIR pyridine adsorption

The nature and strength of acid sites were evaluated by infrared spectroscopy using pyridine as a probe molecule. The FTIR spectra are shown in Figure 4. The adsorption of pyridine on Lewis acid sites was observed in all catalysts, as evidenced by the bands at 1444cm$^{-1}$, 1490cm$^{-1}$ and 1600cm$^{-1}$ (Manríquez et al., 2004). No band near 1540cm$^{-1}$, assigned to pyridine adsorbed on Brönsted sites, was clearly observed.

FTIR spectra confirmed that pyridine was not completely desorbed at 350°C for TiO$_2$-P and Nb$_2$O$_5$-P, which indicates the existence of stronger Lewis sites when compared to untreated catalysts. Table 3 shows the amount and density of acid sites. It also provides a classification of weak, moderate, and strong acid sites based on the fraction of sites occupied by pyridine after desorption prior to reaching 150°C, between 150 and 250°C, and above 350°C, respectively.
The amount of acid sites of titanium catalyst had a significant increase after the H₃PO₄ treatment, while the niobium catalyst showed a reduction of sites. This reduction has already been reported by Okasaky and Kurimata (1990). According to the authors, the decrease in the amount of acid sites is attributed to the size of pyridine molecules, which are excessively large to penetrate in the micro-pores of the surface layer formed by the phosphoric acid treatment. This phenomenon was not observed in the titanium catalyst, probably due to a different surface layer structure with larger pores formed after phosphoric acid treatment.

This non-volatile layer of polyphosphate over the catalyst surface is said to be responsible for the increase of the amount of catalyst acid sites (Okazaky et al., 1987).

The increase in the amount of acid sites could not be observed after acid the treatment over niobium catalyst. Therefore, the pronounced effect on the total density of moderate and strong acid sites on this catalyst, leading to an increase from 42.5 to 60.4% after treatment, confirms the

**Table 3. Acid properties of the catalysts.**

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>Amount of acid sites (μmol_pyridine/g)</th>
<th>Density of acid sites (μmol_pyridine/m²)</th>
<th>Weak acid sites (%)</th>
<th>Moderate acid sites (%)</th>
<th>Strong acid sites (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>TiO₂</td>
<td>16</td>
<td>2.0</td>
<td>77.0</td>
<td>23.0</td>
<td>0.0</td>
</tr>
<tr>
<td>TiO₂-P</td>
<td>90</td>
<td>10.6</td>
<td>91.0</td>
<td>0.8</td>
<td>8.2</td>
</tr>
<tr>
<td>Nb₂O₅</td>
<td>65</td>
<td>0.78</td>
<td>57.5</td>
<td>42.5</td>
<td>0.0</td>
</tr>
<tr>
<td>Nb₂O₅-P</td>
<td>49</td>
<td>0.67</td>
<td>39.6</td>
<td>27.0</td>
<td>33.4</td>
</tr>
</tbody>
</table>
effectiveness of phosphoric acid treatment in promoting the increase of catalysts acid strength. Likewise, this effect was observed by Tang et al. (2010). In this work, the authors attributed the increase of acid strength to the formation of POH acid site on catalyst surface, which are stronger than the NbOH sites.

### 3.5 Glucose dehydration

Table 4 presents the initial reaction rates of glucose consumption, fructose and HMF formation at 200°C.

Regarding glucose consumption and HMF formation, there is a clear and direct relation between density of acid sites and initial reaction rate. However, it is relevant to note that, in the case of titanium catalysts, where the initial rates for glucose consumption and HMF formation are more important, although the phosphated catalyst showed a 5.6 times higher density of acid sites (see Table 3) as compared to the untreated titanium catalyst, the initial rates for glucose consumption and HMF formation were only 1.3 and 1.1 times higher than the untreated catalyst, respectively. These results suggest that the relationship between density of moderate/strong and weak acid sites, higher for the untreated titanium catalyst, is more appropriate for HMF formation.

As mentioned, fructose formation via glucose isomerization can be promoted either by proton transfer, by a basic-catalyzed reaction, or by hydride transfer catalyzed by Lewis acid centers. However, large amounts of catalysts with Lewis sites are required to perform glucose isomerization (Nakajima et al., 2011). Since a catalyst/glucose mass ratio of only 0.10 was used in this work, the hydride transfer mechanism was not expected to be as important as the proton transfer mechanism.

In Table 4 it can be seen that, with titanium catalysts, the initial reaction rates for fructose formation are around 5-10 times higher when compared to the niobium catalysts. However, the first ones also have a much higher density of basic sites than the latter ones (Table 2), which could explain the significant difference between initial rates.

Although the untreated titanium catalyst showed a higher initial rate for fructose formation, it had almost half of the density of basic sites observed in the H₃PO₄ treated catalyst. However, the untreated catalyst showed weaker basic sites when compared to the treated catalyst by CO₂-TPD, suggesting that weak basic sites are more important than strong basic sites for fructose formation from glucose isomerization. This assertion is in agreement with the literature (Watanabe et al., 2005B). Comparing the niobium catalysts among themselves, a slightly-higher fructose initial rate was observed with Nb₂O₅-P, which also had a higher density and weaker basic sites than Nb₂O₅.

Figure 5 shows glucose conversions and HMF yields as a function of residence time for all tested catalyst at 200°C.

HMF yields follow a bell-shaped profile, usually observed in consecutive reaction systems. Indeed, the HMF formed is subsequently decomposed into organic acids and humins, as mentioned before. For all catalytic tests, the formation of humins increased with reaction time. For longer periods of reaction the formation of this oligomeric material caused severe catalyst deactivation, covering the entire catalyst surface and granting a muddy aspect to the reaction media. Nevertheless, Figures 5A and 5B indicate catalytic activity for both glucose consumption and HMF decomposition at longer reaction times, probably due the organic acids formed from HMF.
Trends for fructose and HMF yields and glucose conversion for each catalyst are shown in Figure 6. At lower glucose conversions (below 60% mol/mol), it was observed that catalysts with weaker basic sites had higher fructose yields, corroborating to what was observed when analyzing fructose initial rates. However, most likely due to the higher amount of moderate/strong acid sites, faster glucose consumption was noticed for niobium catalyst, leading to an abrupt reduction in fructose yield. Although the titanium catalysts had shown lower fructose yields at lower glucose conversions, the presence of considerably lower amount of moderate/strong acid sites allowed gradual fructose consumption.

At higher conversions (above 70% mol/mol), a process condition where the deactivation by humins formation is more severe, the niobium catalysts showed lower HMF yields than the titanium catalysts.

These results could be associated with the higher fructose availability at higher glucose conversion for titanium catalysts, and with the considerably higher amount of moderate/strong acid sites presented in the reaction medium when niobium catalysts were used. Although these strong acid sites promote glucose and fructose consumption and the HMF formation, they also promote the decomposition of HMF in organic acids and humins.

Among all catalysts, the H$_3$PO$_4$ treated titanium catalyst achieved the highest HMF at high glucose conversions. High densities of basic sites generated after acid treatment associated to its low acid strength probably allowed this solid to keep its
activity for longer periods and to be more selective than the other catalysts. This result could be even more pronounced if its basic sites strength could be reduced with the phosphation processes.

4. CONCLUSIONS

The effects of $\text{H}_3\text{PO}_4$ treatment on the catalytic activity of niobium and titanium oxides were evaluated on glucose dehydration in aqueous medium. The activities under study were related to catalyst acid and basic properties. The experiments confirmed that weak basic sites favor glucose isomerization into fructose while catalysts with moderate and strong acid sites favor not only the formation of HMF and glucose conversion, but also HMF decomposition in organic acids and humins. Therefore, in order to increase the HMF yield, a compromise between moderate acidity, weak basicity and high basic and acid sites densities must be achieved.

Thus, phosphated TiO$_2$ stands out as a more promising catalyst for glucose dehydration. However, the severe deactivation process observed for all catalysts indicates that stability and regeneration tests must be carried out to develop guidelines for the continuous process for synthesis of HMF from glucose in aqueous medium.

Figure 6. Fructose yield versus glucose conversion (A) and HMF yield versus glucose conversion (B) at 200°C.
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