Selectivity Control during the Single-Step Conversion of Aliphatic Carboxylic Acids to Linear Olefins

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ABSTRACT: We have studied the single-step catalytic conversion of biomass-derived aliphatic carboxylic acids to linear olefins via tandem hydrogenation/dehydration reactions. Hexanoic acid was converted to a mixture of hexenes (92.0% selectivity) over silica–alumina supported Cu nanoparticles. Remarkably, we observed a rapid selectivity switch to 99.8% hexane once carboxylic acids were fully consumed, with similar results using butanoic acid derived from biomass using consolidated bioprocessing. Based on intermediate, desorption, and in situ spectroscopy studies, we propose that the presence of a small amount of carboxylic acid on the catalyst surface leads to a dramatic decrease in overhydrogenation of olefins.

KEYWORDS: olefins, carboxylic acids, copper, silica–alumina, FTIR spectroscopy, selectivity control, biomass

Utilization of lignocellulosic biomass is becoming increasingly important as we attempt to reduce our dependence on fossil fuels, which are increasingly unsustainable and environmentally problematic. In addition to being used as a source of energy, we are especially reliant on using fossil fuels as our main carbon source for producing carbon-based chemicals and other industrial compounds. Linear olefins, a major bulk chemical and a precursor for, among others, plastics and detergents, are an example of a major product that is derived from petroleum.

Biomass-derived carboxylic acids could be a potential replacement for petroleum in the production of olefins. Carboxylic acids are abundant in nature, natively present in vegetable oil as triglycerides, with more than 350 plants known to contain oil. Alternatively, lignocellulosic biomass can be converted to carboxylic acids via fermentation, either by using mixtures of simple sugars derived from biomass or directly fermenting pretreated biomass, resulting in aliphatic carboxylic acids of 2 to 6 carbon length depending on the fermentation process.

Deoxygenation of aliphatic carboxylic acids to olefins has always been a challenge because of the high stability of the carboxyl group and the lack of any electron-withdrawing groups (e.g., phenyl groups) or neighboring hydroxyl functionalities (α-hydroxy acids). Aliphatic carboxylic acids are typically weak acids with low dissociation constants (pK_a of hexanoic acid = 4.85). Furthermore, dissociated carboxylate anions are resonance-stabilized, thus lowering their reactivity. The straight chain carbon backbone also limits any reactions to carbon–carbon bond scissions. Thus, the upgrading process must start by targeting the carboxyl group.

Traditional thermal decarboxylation processes usually result in alkanes rather than olefins. Oxidative decarboxylation is a possible route to produce olefins, with the loss of one molecule of carbon dioxide. Decarboxylation of carboxylic acids, where a carbon monoxide and water molecules are removed leaving behind an olefin, has also been demonstrated. However, both routes require the use of a stoichiometric oxidant/reagent to function, which is unsustainable and expensive in the context of bulk chemical production. Environmental friendly oxidants such as H_2O_2 or O_2 can be used in enzymatic oxidative decarboxylation, but these systems use specifically engineered strains of bacteria and also require additional cofactors.

Rather than decarboxylation, it is possible to perform hydrogenation. This maintains the carbon chain length, which is especially important in the context of biomass conversion due to the importance of maximizing carbon conversion. However, care must be taken not to fully hydrogenate the carboxylic acid to an alkane. Platinum or copper can be used to hydrogenate carboxylic acids to...
alcohols. The alcohol formed could then be dehydrated to olefins over an acid catalyst, such as sulfuric acid or silica-alumina.

In this work, our objective was to devise an alternative process to convert dilute streams of carboxylic acids to olefins in one step using tandem hydrogenation/dehydration. By first hydrogenating the carboxylic acids to alcohols, followed by subsequent dehydration to olefins in the same reactor, we can avoid expensive sacrificial reagents as well as preserve the carbon chain length of the substrate. In our investigation of using hexanoic acid and butanoic acid as substrates, we discovered a sudden switch in olefin/alkane selectivity once carboxylic acids were fully consumed. Using several catalyst surface studies, we were able to determine the nature of this selectivity switch. Finally, we demonstrate the application of this process to butanoic acid that was directly produced from pretreated beech wood in a consolidated bioprocess.

Performing tandem hydrogenation/dehydration reactions on carboxylic acids in one process required a bifunctional catalyst that had both hydrogenation and alcohol dehydration properties. We selected Cu supported on commercial silica-alumina (Cu/Si–Al), with the rationale that Cu could act as a mild hydrogenation catalyst and the acid sites on silica-alumina could perform dehydration reactions. Partial hydrogenation of hexanoic acid resulted in 1-hexanol, which could partially esterify with unreacted hexanoic acid to form hexyl hexanoate (Scheme 1). This is an undesired side reaction that occurs mainly on Brønsted acid sites, but it is reversible.21 As an intermediate, 1-hexanol then underwent dehydration to form hexenes. As a primary alcohol, the reaction likely occurred through the E2 mechanism, resulting in 1-hexene. However, at high temperatures and acidic conditions, double-bond migration was favorable, resulting in formation of internal hexenes. Finally, undesired overhydrogenation of the hexenes to hexane also occurred, and, as discussed below, was highly dependent on reaction conditions.

We carried out the tandem hydrogenation/dehydration of hexanoic acid in an upflow fixed-bed reactor at 210 °C and 5 bar H₂ using Cu/Si–Al with a copper loading of 6 wt % and a metal dispersion of 5%. At first, hexane was the major product indicating almost complete overhydrogenation of olefins (Figure 1A), likely due to an abundance of hydrogen present on the fresh catalyst bed. After this initial induction period, the reaction stabilized and selectivity to hexenes increased significantly until they were the major product. At a weight hourly space velocity (WHSV) of 0.99 h⁻¹, we obtained 96.7% conversion and a molar product distribution of 92.0% hexenes and 7.3% hexane at steady state. The hexenes consisted of predominantly 2-hexene and 3-hexene (95 mol % of total olefins), indicating a high degree of double bond migration after the dehydration step. Skeletal rearrangement was not significant (3 mol % of total olefins). There was a slight overhydrogenation of the olefins to hexane under those reaction conditions, along with a small production of hexyl hexanoate. The conversion and product distribution were measured using a GC-FID and a GC-MS. The conversion and product distribution were calculated based on the total liquid feed.

Scheme 1. Proposed Reaction Pathway for the Tandem Hydrogenation/Dehydration of Hexanoic Acid.24

Figure 1. (A) Conversion and molar product distribution of hexanoic acid tandem hydrogenation/dehydration as a function of time on stream (WHSV = 0.99 h⁻¹, H₂ flow = 74 mL/min, T = 210 °C, P = 5 bar and feed = 2 wt % hexanoic acid in isooctane). (B) Average steady state conversion and molar product distribution of hexanoic acid tandem hydrogenation/dehydration at different WHSV (T = 210 °C, P = 5 bar and feed = 2 wt % hexanoic acid in isooctane). (C) Molar product distribution of an intermediate products study as function of time on stream (WHSV = 0.99 h⁻¹, H₂ flow = 74 mL/min, T = 210 °C, P = 5 bar and initial feed = 2 wt % 1-hexene in isooctane). Green lines represent a feed switch to either 2 wt % 1-hexene/2 wt % hexanoic acid in isooctane or 2 wt % 1-hexene in isooctane. The legend for Figure 1B,C can be found in Figure 1A. The WHSV is calculated based on the total liquid feed.

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constant over 80 h on stream (0.8 g hexenes produced/g catalyst over 80 h in this experiment), attesting good catalyst stability. When using a higher concentration (10 wt %) of hexanoic acid as feed (and adjusting the flow rate and WHSV to yield a similar ratio of incoming hexanoic acid to catalyst weight), a comparable performance to the 2 wt % feed was achieved, demonstrating that a high dilution was not necessary to obtain these results (Figure S1). Increasing the WHSV to 1.23 h⁻¹ lowered the conversion to 87.5% (Figure 1B). At these conditions, we also detected a higher amount of hexyl hexanoate in the product stream, showing that esterification was a competing side reaction when significant amounts of hexanoic acid were present. In the previous case of 96.7% conversion, the amount of unreacted hexanoic acid was low enough that most of the 1-hexanol formed proceeded to dehydrate to hexenes. We lowered the temperature of the reaction to 200 °C, thereby further lowering the conversion to 77.8%. At this conversion, large quantities of hexyl hexanoate were formed. The proportion of hexane formed was similar compared to previous conditions, implying that the slight overhydrogenation of olefins was independent of the conversion and WHSV.

Interestingly, we observed a brusque selectivity switch to 99.8% hexane at full conversion, after a decrease in WHSV from 0.99 h⁻¹ to 0.74 h⁻¹ (while maintaining the hydrogen flow to feed flow ratio constant). This contrasted with other reaction conditions where the proportion of overhydrogenation was constant.

We hypothesize that the presence of a small amount of hexanoic acid on the catalyst surface inhibited overhydrogenation of the hexenes to hexane. To ascertain the nature of this selectivity switch, we performed a study of intermediate products at WHSV = 0.99 h⁻¹ (Figure 1C). By initially flowing only 1-hexene over the catalyst bed, complete overhydrogenation was observed. Changing the feed to a blend of hexanoic acid and 1-hexene switched the selectivity to a similar product distribution as the original hexanoic acid feed (88.3% hexenes and 11.2% hexanes) at those conditions. As the reaction temperature and pressure were constant, this switch could not have happened because of changes in vapor pressure or an unexpected phase change. We rationalized that this phenomenon occurred as a result of the interaction of the reaction species on the surface of the catalyst in the presence/absence of hexanoic acid.

The initial induction period of the reaction (shown for a WHSV of 0.99 h⁻¹ in Figure 1A) was also used to probe this phenomenon. When conducting the experiment under Ar flow for the first 24 h (instead of H₂) and subsequently switching to H₂ flow, a high selectivity of hexenes was achieved immediately (Figure S2A). However, this resulted in some deactivation of the catalyst. When performing a similar experiment using an Ar flow for 1 h (Figure S2B), the deactivation was not as significant (83.4% vs 96.7% conversion), and a shorter induction period was observed compared to constant H₂ flow (7 h vs 24 h). These results showed that H₂ was important in maintaining catalyst stability and confirmed that the initial induction period with high alkane selectivity was due to an abundance of hydrogen present on the surface of a fresh catalyst bed. Furthermore, preadsorbing the catalyst with hexanoic acid mitigated the initial overhydrogenation of the hexenes to hexane, albeit with severe catalyst deactivation under an Ar atmosphere, further confirming the important effect of adsorbed carboxylic acids on product selectivity.

To further explore the interaction of the reacting species with the catalyst surface, a temperature-programmed desorption (TPD) of 1-hexene on the catalyst surface was performed to determine its adsorbed quantity on the catalyst (Figure 2A).

![Figure 2](image)

On a clean catalyst surface, we observed 234 µmol/g of 1-hexene adsorbed on the surface of the catalyst. In contrast, by preadsorbing hexanoic acid on the catalyst (which, we estimate led to roughly the same amount of acid adsorbed as 1-hexene—measured as 285 µmol/g with acetic acid, see Section S of the SI), the amount of 1-hexene adsorbed decreased to 10 µmol/g. TPD of just hexanoic acid confirmed the attribution of the 2 desorption peaks at ~400 °C and ~700 °C to the desorption and degradation of hexanoic acid itself (the fragmentation pattern of hexanoic acid contains mass 84), and not to more strongly adsorbed 1-hexene in the presence of hexanoic acid. These findings suggest that the presence of hexanoic acid severely limits the adsorption of 1-hexene, thus decreasing overhydrogenation.

We further confirmed this hypothesis by Fourier-transform infrared (FTIR) spectroscopy. 1-Hexene adsorbed on a clean catalyst exhibited two adsorption bands at 1608 and 1449 cm⁻¹, attributed to the C=O stretching mode and the CH₂ scissoring mode, respectively (Figure 2B). The slight red-shift of the C=O band is attributed to C=C bond weakening upon adsorption on the catalyst. Conversely, the characteristic bands of 1-hexene were not observed upon 1-hexene adsorption.
adsorption after preadsorption of hexanoic acid on the catalyst. The absence of 1-hexene signals further indicated that hexanoic acid prevented 1-hexene from bonding and being overhydrogenated to hexane by occupying the hydrogenation sites of the catalyst (Figure 2C). This phenomenon can be extended to other carboxylic acids. We performed tandem hydrogenation/dehydration on butanoic acid and obtained an average steady-state molar product distribution of 90.9% butenes, 3.7% butane, and 5.4% others at a conversion of 90.9% (Figure 3A). The increased amount of esters can be due to the lower conversion of 90.9% compared with the 96.7% obtained with our previous experiments using hexanoic acid. Nonetheless, by pushing the conversion to completion, we observed a similar selectivity switch, although there was a higher proportion of butenes (17.0%) to butane (83.0%) compared with hexanoic acid (99.8% hexane). The feed flow rate had to be reduced for butanoic acid, as there was a higher molar concentration in the feed due to the lower molar mass of butanoic acid compared with hexanoic acid.

In order to demonstrate this concept for real biomass-derived products, we used our process on butanoic acid produced with a novel consolidated bioprocessing method that produced the carboxylic acid directly from steam-exploded beech wood (Figure 3B). Tandem hydrogenation/dehydration of biomass-derived butanoic acid initially proceeded similarly to commercial butanoic acid feed, with a peak molar product distribution of 80% butenes (Figure 3C). However, the catalyst suffered from significant deactivation after 8 h on stream. It is likely that the extraction process resulted in slight impurities that adsorbed on the catalyst surface, as the reaction proceeded smoothly with the commercial butanoic acid feed. Nevertheless, this shows that the use of biomass-derived carboxylic acids was possible, although optimization of the extraction process is still required.

In summary, we demonstrate that biomass-derived streams of carboxylic acids can be converted into mixtures of olefins in a single step. Using Cu/SiO2−Al2O3, > 90% olefin selectivity was achieved at close to 99% conversion. At full conversion, an abrupt selectivity switch occurred, changing the product distribution from predominantly olefins to almost exclusively alkanes. Through TPD and FTIR catalyst surface studies, we proposed that the observed selectivity switch occurred because of the presence of a small amount of carboxylic acids on the catalyst surface, which prevented binding and subsequent overhydrogenation of the olefins. We propose that the inhibition of catalyst active sites using carboxylic acids could also be applied to hydrogenation reactions using other substrates, where selective preservation of double bond functionalities is required. By doing so, we could possibly influence the product distributions of other types of catalytic reactions by controlling the presence of carboxylic acids in the reaction media. Differences in catalyst performance when processing commercial compounds versus real biomass-derived feed in biomass upgrading reactions could also be explained; especially given the ubiquity of carboxylic acid impurities in biomass-derived mixtures. An improved understanding of the influence of these functionalities on catalytic reactions will be important in developing future biorefining processes. Therefore, this phenomenon can both help better understand the
catalytic processing of biomass and open new avenues for selectivity control during catalytic reactions.

**REFERENCES**


