Kinetics of the Homogeneous Catalytic Hydrogenation of Olefins in Supercritical Carbon Dioxide Using a Fluoroacylate Copolymer Grafted Rhodium Catalyst

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The kinetic studies on the hydrogenation of 1-octene and cyclohexene using a fluoroacylate copolymer grafted rhodium catalyst in supercritical carbon dioxide (scCO2) are reported. The reactions were investigated at temperatures between 50 and 120 °C, and pressures ranging from 172 to 241 bar. The catalyst also deactivated at these reaction conditions. For the case of 1-octene, isomerization to (E)2-octene and (Z)2-octene also occurred as side reactions. To represent the experimental data, a kinetic model was developed on the basis of reported studies about hydrogenation and isomerization of olefins in conventional solvents. It was proposed that two hydride catalytic species are formed during the kinetic cycle. Monohydride species promoted the isomerization, and dihydride species catalyzed the hydrogenation. Deactivation of the catalyst was attributed to the formation of an unsaturated olefin complex. Statistical methods were applied to discriminate among rival models. It was found that the rate-determining steps for hydrogenation and isomerization were the formation of the alkylrhodium hydride complex and the coordination of the olefin in the monohydride catalytic species, respectively.

Introduction

Many traditional organic solvents employed in homogeneous catalysis, such as benzene, toluene, methylene chloride, THF, and DMF, are being phased out as a result of U.S. FDA and U.S. EPA regulations to limit emissions of and exposures to potentially toxic and environmentally damaging compounds. Moreover, in homogeneous catalytic processes, other disadvantages include the expensive catalyst recovery and purification processes after the reaction, and the low solubility of gases in liquid solvents.1–2 Supercritical fluids (SCFs) have shown significant potential as an alternative to replace the common organic solvents as reaction media because of their unique physicochemical properties. Several reviews have been published explaining the fundamentals and the main advances in this new field of chemical reaction engineering.3–8 Specifically, supercritical carbon dioxide (scCO2) is an excellent alternative to conventional solvents because it is easily separated from the other components in the process stream, nonflammable, safe to human exposure, acceptable for environmental release, and capable of performing the duties of a nonpolar solvent while allowing manipulation of its solvent strength through a wide range of polarities. At supercritical conditions, its density is sufficient to provide dissolution strength, the diffusivity of solutes is higher than in liquids, and the viscosity is lower increasing the mass transfer; therefore, a hydrogenation reaction that is diffusion-limited in the solvent phase can be enhanced by carrying it out in an SCF due to higher diffusivity and elimination of gas/liquid and fluid/fluid interphases. The main drawback of using scCO2 as reaction media is the scarce solubility of conventional organometallic catalysts. Therefore, efforts have been focused on the design of new catalytic materials that are miscible in scCO2 under mild conditions. Particularly, fluoropolymers and silicones have moderate solubility in scCO2.9 On the basis of that, new catalysts have been developed by grafting the catalytically important ligands to a “CO2-philic” backbone.10–14 Catalytically active sites become available in scCO2 by the fixation of “CO2-philic” solubilizers in the periphery of the ligand.

In the present work, a novel Rh catalyst, soluble in scCO2, was synthesized by attaching rhodium ligands to a fluoroacrylate copolymer. Kinetic studies on the hydrogenation of 1-octene and cyclohexene with this novel catalyst were made. It was found that the catalyst deactivated at reaction conditions. Also, 1-octene was isomerized to (E)2-octene and (Z)2-octene as side reactions, and no appreciable hydrogenation of the 2-octene isomers was detected. Finally, the kinetic model for the hydrogenation and isomerization processes was established.

Experimental Section

Materials. Organic substances were purchased from Sigma-Aldrich and used as received. Carbon dioxide and hydrogen were bought from Brazos Welding Supply and used as received. Catalyst was prepared in the Department of Chemistry at Texas A&M University and used as received.

Catalyst. The catalyst employed in the experiments was a Wilkinson-type catalyst, where a rhodium organometallic complex was grafted to a polymer soluble in scCO2. Details about the synthesis of the catalyst have been given elsewhere.15 First, the polymer was synthesized by the polymerization of 1H,1H,2H,2H-heptade-
cafluorodecyl acrylate monomer (zonyl TAN) and N-acrylosuccinimide (NASI), the former increasing the solubility in scCO₂ and the latter providing attachment sites for the catalyst. ¹H NMR analysis confirmed the zonyl TAN to NASI molar ratio, which was adjusted between 7 and 20. Then, the succinimide group was exchange by phosphines using NH₂(CH₂)₃PPh₂ (DPPA). During this step, 15–30% of the phosphines were oxidized due to uncontrolled reaction as was estimated by ³¹P NMR analysis. Finally, this precursor was reacted with [RhCl(COD)]₂ to obtain the catalyst. Neutron activation analysis gave a catalyst composition of 10⁻⁸ mol of Rh per gram of catalyst. The ideal molecular structure of the catalyst is presented in Figure 1.

**Apparatus and Procedures.** The experimental setup and procedures of operation have been described elsewhere. In short, the system consisted of a 100 mL EZE-seal vessel with a magnetic drive stirrer by Autoclave Engineers and warmed with a heating jacket. The temperature was regulated using a temperature controller (Omega CN9000A) and monitored with a thermocouple (Omega 115KC) located in a thermowell inside the reactor. The pressure was measured using an Autoclave gauge. As a safety measure, a rupture disk was varied according to the olefin concentration. The amount of catalyst added to the reactor was 0.53 mm ID × 1.50 mm OD × 30 m, the latter providing attachment sites for the catalyst. ¹H NMR analysis confirmed the zonyl TAN to NASI molar ratio, which was adjusted between 7 and 20. Then, the succinimide group was exchange by phosphines using NH₂(CH₂)₃PPh₂ (DPPA). During this step, 15–30% of the phosphines were oxidized due to uncontrolled reaction as was estimated by ³¹P NMR analysis. Finally, this precursor was reacted with [RhCl(COD)]₂ to obtain the catalyst. Neutron activation analysis gave a catalyst composition of 10⁻⁸ mol of Rh per gram of catalyst. The ideal molecular structure of the catalyst is presented in Figure 1.

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**Figure 1.** Ideal molecular structure of the catalyst.

After each experimental run, the reactor was washed with acetone to remove any hydrocarbon residue, 1,1,2-trichlorotrifluoroethane to dissolve the remained catalyst, and nitric acid to eliminate any metallic catalytic deposit that could be formed during the experiment. Finally, a mechanical cleaning was performed in all the metallic parts.

Then, a blank reaction test was performed to verify total removal of the catalyst. If conversion was obtained during the blank test, the cleaning procedure was repeated until no conversion was observed.

Samples taken from the reactor were analyzed by gas chromatography (HP 5890 equipped with a flame ionization detector) using a capillary ZB-5 column (5% Phenyl Polysiloxane, 30 m × 0.53 mm ID × 1.50 μm FT) by Phenomenex.

**Hydrogenation Reaction Model.** The hydrogenations of 1-octene and cyclohexene were selected to determine the activity of the catalyst. These reactions are exothermic and progress at low temperatures to reach total conversion without thermodynamic limitations. Isomerization to 2-octene and 3-octene may also occur with the Wilkinson-type catalysts. Isomerization reactions are also exothermic proceeding at mild temperatures. The initial mole fractions of all components were kept constant at all temperatures; this means that, at lower temperatures or higher pressures, where the density of scCO₂ is higher, the concentrations of hydrogen and olefin were greater. For the hydrogenation of 1-octene, the initial mole fractions of olefin and hydrogen were 0.0010 and 0.0334, respectively. In the case of hydrogenation of cyclohexene, the initial mole fractions of olefin and hydrogen were 0.0002 and 0.056, respectively. The amount of catalyst added to the reactor was varied according to the olefin concentration. The temperature and pressure range studied were 50–95 °C and 172–241 bar. Experiments were run twice for reproducibility.

**Analysis and Discussion**

Estimations of the theoretical critical point of the reaction mixtures as the reactions proceeded were performed using the group contribution method for the critical temperature, and the method of Liu for the critical pressure. It was found that in both cases the critical temperature slightly changes from 301.47 to 301.54 K, and the critical pressure varies from 83.18 to 83.00 bar. Therefore, it can be concluded that reactions always proceed under supercritical phase conditions.

Preliminary experiments at 172 bar and different temperatures were done to determine the best substrate to catalyst molar ratio to perform parameter estimation for the hydrogenation of 1-octene. The catalyst employed was RhCl(TAN₁₅DPPA)₂ with 15% of oxidized phosphines. Results are presented in Figures 3 and 4. When the substrate to catalyst molar ratio was 800, excellent reaction profiles were obtained at 70 and 95 °C; however, at 50 °C almost no conversion was obtained. On the other hand, when this molar ratio decreased to 400 very fast conversion was observed at 70 and 95 °C, but fair reaction profile was observed at 50 °C. Therefore, the selected substrate to catalyst molar ratio was 400 for experiments at 50 °C, and 800 for experiments at 70 and 95 °C.

**Kinetic Mechanism.** The hydrogenation of unsaturated compounds using organometallic catalyst takes place through several steps involving oxidative-addition, coordination, insertion, and reductive-elimination reactions. The hydrogenation mechanism goes through intermediate metal hydrides. The formation of metal hydrides typically takes place by oxidative-addition. Monometalic and bimetallic H₂ oxidative-additions are known, and both need the presence of at least one vacant site on the metal as is presented in Figure 5. Many factors influence the reactivities of metal complexes to achieve oxidative-additions. Electron-releasing ligands, i.e., phosphines, enhance the tendency to undergo oxidative-additions; in contrast, π-acceptors, such as CO or olefins, decrease the propensity to accomplish oxidative-additions. Additionally, steric inhibition is also significant, particularly with large phosphine groups. Oxidative-addition of C=C bonds is also thermodynamically possible since the energetics of C=C oxidative additions are sometimes not much different from H₂ additions. Thus,
the olefin is coordinated forming an unsaturated olefin complex as is presented in Figure 6.

Once the oxidative-addition has been accomplished, these new species either coordinate an unsaturated hydrocarbon, or add H₂. However, it has been suggested that the formation of hydride olefin complexes from an unsaturated olefin complex is difficult, and in fact, the formation of an unsaturated olefin complex may tend

Figure 2. Schematic diagram of the reactor system.

Figure 3. Preliminary experiments for the hydrogenation of 1-octene at 172 bar using a substrate to catalyst molar ratio of 800. Temperature: (●) 50 °C, (■) 70 °C, (▲) 95 °C.

Figure 4. Preliminary experiments for the hydrogenation of 1-octene at 172 bar using a substrate to catalyst molar ratio of 400. Temperature: (●) 50 °C, (■) 70 °C, (▲) 95 °C.

to inhibit the hydrogenation process. Therefore, it is considered that only the mono- and dihydride species may continue promoting the hydrogenation and isomerization.

Homogeneous hydrogenation catalysts with dihydride complexes are much better studied and established. A generalized mechanism is given in Figure 7. Coordination of an olefin on the same face of the coordination sphere as the two hydrides provides the coordinative saturated complex 5. The transformation of 2 to 5 requires a vacant coordination site, accounting for some of the inhibition of the catalytic cycle brought on by competing extraneous ligands. The migratory insertion of 5 to 6 goes by cis addition of the M-H bond to the olefin and requires the olefinic bond axis, the metal, and the hydride to be coplanar. This step is usually assisted or consummated by the incorporation of the external ligand. Alkyl-hydride intermediates such as 6 have rarely been observed because the following step is very rapid, preventing buildup of 6 giving high stereoselectivity and the absence of olefin rearrangement.

From the preceding explanation, it could be concluded that dihydride catalytic species would not produce
isomers; therefore, a different catalytic species should be generated to produce isomers. This species could be a rhodium monohydride complex, which is an excellent isomerization catalyst. The formation of this complex is uncertain, but it has been proposed that it may proceed through heterolytic dihydrogen activation to finally obtain a monohydride RhI catalytic species. For simplification in the deduction of the kinetic mechanism, it could be assumed that the generation of this complex proceeds in a single step as it was presented in Figure 5b. Once the monohydride species has been formed, the next step involves complexation of the olefin to a coordinatively unsaturated hydride, followed by a Markovnikov hydride migration, complex 8, as is presented in Figure 8. Then, \( \gamma - H \) (not \( \alpha - H \)) elimination would take place to produce an internal olefin as is depicted in Figure 8.

Finally, it was considered that the catalyst was inhibited due to the formation of an unsaturated olefin complex as it was presented in Figure 6. The way this complex is formed is uncertain; it could be either by oxidative-addition to form a metallacyclopropane-type complex, or by simple coordination of the double bond to the metal. For simplicity, the last reaction was assumed. This complex could further react to produce dimer species that could be precipitated during the reaction.

Figure 9 presents the proposed reaction mechanism for the production of n-octane and 2-octene isomers. Here, the cycle for the generation of isomer must be repeated twice with different rate constants, one for the production of (E)2-octene, and the other for the production of (Z)2-octene; this is the reason that two reaction constants appear in the different reaction steps.

**Parameter Estimation.** For determining the kinetic model for the hydrogenation of 1-octene, models were proposed assuming different combinations of rate-determining steps for hydrogenation and isomerization reactions as is summarized in Table 1. Because hydrogen was fed with huge excess with respect to olefin, steps 1 for hydrogenation and 5 for isomerization were discarded as rate-controlling steps. Moreover, considering any of these steps as rate-controlling would lead to obtaining a kinetic expression just in terms of hydrogen concentration (without effect of the olefin concentration), and since hydrogen is in great excess, this kinetic expression would be meaningless.

<table>
<thead>
<tr>
<th>model</th>
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<th>isomerization</th>
</tr>
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<tbody>
<tr>
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</tr>
<tr>
<td>2</td>
<td>3</td>
<td>9, 13</td>
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<tr>
<td>11</td>
<td>3</td>
<td>6, 10</td>
</tr>
<tr>
<td>12</td>
<td>2</td>
<td>6, 10</td>
</tr>
</tbody>
</table>
Parameter estimation was performed using Simusolv. It was considered that the equilibrium constants were not affected by pressure at constant temperature, and only the kinetic constant for the rate-determining step is influenced by pressure. Our fits showed that the pressure variation of the equilibrium constants were less than one standard deviation and also there is no change in the number of moles with the reaction. Hence it is assumed that the equilibrium constants are pressure independent. Models 1–4 and 9–10 were rejected because of lack of fit of the experimental data. Also, models 5 and 6 were eliminated because the standard deviations of some computed constants were greater than their numerical value.

The remaining models had the % variation explained (basically regression coefficient) presented in Table 2. Simusolv calculated the % variation explained according to

$$r^2_{overall} = 1 - \frac{\sum_{j=1}^{m} \text{SSQ}_j}{\sum_{j=1}^{m} \text{SSQ}_{\text{mean}j}}$$

and

$$\text{SSQ}_{\text{mean}j} = \sum_{i=1}^{n} \frac{(y_{ij} - \bar{y}_j)^2}{f_{i,j}^\gamma}$$

$$\text{SSQ}_j = \sum_{i=1}^{n} \frac{(y_{ij} - f_{i,j})^2}{f_{i,j}^\gamma}$$

with

$$\bar{y}_j = \frac{\sum_{i=1}^{n} y_{ij}}{\sum_{i=1}^{n} f_{i,j}^{\gamma/2}}$$

$$\gamma = \frac{\sum_{i=1}^{n} (y_{ij} - \bar{y}_j)^2}{\sum_{i=1}^{n} f_{i,j}^{\gamma/2}}$$

where $n$ is the number of observations, $m$ is the number of variables, $y$ is the experimental value, $f$ is the predicted value, and $\gamma$ is a weighing parameter.

From Table 2, no discrimination could be done; therefore, statistical methods were employed to discriminate among rival models.

First, the likelihood ratio as a discrimination criterion was used. The ratio of the maximum values of the likelihood function of both models was computed according to the following:

$$\frac{(L_1)_{\text{max}}}{(L_2)_{\text{max}}} = \left( \frac{n - p_2}{n - p_1} \right)^{n/2} \exp \left( \frac{p_1 - p_2}{2} \right) \left( \frac{e_1^T e_2}{e_1 e_1} \right)^{n/2}$$
where $e$ is the vector of residuals $y - \hat{y}$, $n$ the number of experimental data, and $p_i$ is number of parameters in model $i$.

For achieving discrimination between models, two nonlinear models must be made:

$$L_1 \max \begin{array}{l} \leq B \quad \text{model 1 is rejected} \\ \geq A \quad \text{model 1 is preferred} \\ A < \frac{L_{1\max}}{L_{2\max}} < B \quad \text{no decision can be made} \end{array}$$

The model with the smallest residual sum of squares is taken as a reference, and the others models are compared to this reference. The residual sum of squares for the each model is given in Table 3. Model 11 had the smallest residual sum of squares, so the other models were compared with respect to it.

The following values were obtained:

$$\frac{(L_{1\max})}{(L_{2\max})} = 2.3 \times 10^7$$

$$\frac{(L_{1\max})}{(L_{9\max})} = 314$$

$$\frac{(L_{11\max})}{(L_{12\max})} = 1.8$$

$A = 19$

$B = 0.0526$

From these results, models 7 and 8 were discarded. However, no distinction can be made between model 11 and 12.

Then, the method of nonintrinsic parameters was applied. This method supposes that a selection between

Table 3. Residual Sum of Squares for the Proposed Models

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<td>9</td>
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<tr>
<td>12</td>
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Table 4. Parameter Values of the Kinetic Model for the Hydrogenation of 1-Octene at 50°C and Substrate to Catalyst Molar Ratio of 400

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<tr>
<th>param</th>
<th>172 bar</th>
<th>207 bar</th>
<th>241 bar</th>
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<td>$4.09 \times 10^{10}$</td>
</tr>
<tr>
<td>$k_6$</td>
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<td>$4.99 \times 10^{10}$</td>
<td>$8.03 \times 10^{10}$</td>
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<tr>
<td>$k_{10}$</td>
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<td>$2.62 \times 10^{10}$</td>
</tr>
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<td>$k_{16}$</td>
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Table 5. Parameter Values of the Kinetic Model for the Hydrogenation of 1-Octene at 70°C and Substrate to Catalyst Molar Ratio of 800

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<td>$4.09 \times 10^{10}$</td>
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<td>$k_6$</td>
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<tr>
<td>$k_{14}$</td>
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<td>$k_{16}$</td>
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<td>$0.20 \times 10^{10}$</td>
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Table 6. Parameter Values of the Kinetic Model for the Hydrogenation of 1-Octene at 95°C and Substrate to Catalyst Molar Ratio of 800

<table>
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<th>param</th>
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<td>$0.20 \times 10^{10}$</td>
<td>$0.14 \times 10^{10}$</td>
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Table 7. Arrhenius Parameter Estimation for the Rate-Determining Step Constants at Given Pressures

<table>
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Table 8. Parameter Values of the Kinetic Model for the Hydrogenation of Cyclohexene Using the Catalyst RhCl(TAN$_3$DPPA)$_3$ with 30% of Oxidized Phosphines at 172 bar

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<th>368 K</th>
<th>393 K</th>
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<td>$8.38 \times 10^{-3}$</td>
<td>$8.38 \times 10^{-3}$</td>
</tr>
</tbody>
</table>
\[ y = \frac{1}{2}(\hat{y}_{11} + \hat{y}_{12}) \]

where \( \hat{y}_i \) is the prediction of the dependent variable \( y \) under model \( i \).

After some mathematical steps, the following is obtained:

\[ E(z) = \lambda(\hat{y}_{11} - \hat{y}_{12}) \]

If model 11 is correct, \( \lambda \) takes a value of \(+0.5\), but if model 12 is correct, its value is \(-0.5\). Regression of \( z \) on \( \lambda(\hat{y}_{11} - \hat{y}_{12}) \) determines which model is best.

Figure 10 presents the plot between \( \lambda(\hat{y}_{11} - \hat{y}_{12}) \) vs. \( z \). Doing linear regression, a value of \( 0.500 \pm 0.0065 \) was calculated for the slope; moreover, Figure 11 shows that the residuals are evenly distributed along the dependent variable confirming the excellent regression analysis. Therefore, model 11 was preferred over model 12.

Thus, if the rate-determining steps are reaction 3 for hydrogenation, and 6 and 10 for isomerization, the following kinetic equations can be derived:

\[
\begin{align*}
\frac{dC_p}{dt} &= \frac{k_3K_1K_2C_AC_BC_{\text{cat}}}{1 + (1 + K_2C_A)K_1C_B + K_6^{1/2}C_{B}^{1/2}} \\
\frac{dC_C}{dt} &= \frac{k_6K_5^{1/2}C_AC_{\text{cat}}C_B^{1/2}}{1 + (1 + K_2C_A)K_1C_B + K_6^{1/2}C_{B}^{1/2}} \\
\frac{dC_D}{dt} &= \frac{k_{10}K_5^{1/2}C_AC_{\text{cat}}B_{\text{cat}}C_B^{1/2}}{1 + (1 + K_2C_A)K_1C_B + K_6^{1/2}C_{B}^{1/2}} \\
\frac{dC_{\text{cat}}}{dt} &= \frac{k_{14}C_AC_{\text{cat}}}{1 + (1 + K_2C_A)K_1C_B + K_6^{1/2}C_{B}^{1/2}} - k_{14}C_{\text{cat}}
\end{align*}
\]

\[ C_A = C_{A0} - C_p - C_C - C_D \]

\[ C_B = C_{B0} - C_p \]

\[ C_{\text{cat}} = C_{\text{cat}0} - C_{\text{cat}} \]

where

\[ A = 1\text{-octene} \]

\[ B = \text{hydrogen} \]

\[ P = n\text{-octane} \]

\[ C = (E)2\text{-octene} \]

\[ D = (Z)2\text{-octene} \]

The values of the parameters are reported in Tables 4–6 for all the studied experimental conditions.

Figures 12–14 show the comparison of calculated and experimental results for the hydrogenation of 1-octene at all the studied experimental conditions. As can be concluded from these figures, the kinetic model was found to be consistent with the general profiles of all the products.

![Graph](image)

**Figure 12.** Comparison between the experimental and calculated results for the hydrogenation of 1-octene at 50 °C: (●) 172 bar, (▲) 207 bar, (▲) 241 bar, (---) calcld.

**Table 9. Arrhenius Parameter Estimation for the Rate-Determining Step Constant of the Hydrogenation of Cyclohexene at 172 bar**

<table>
<thead>
<tr>
<th>constant</th>
<th>( A )</th>
<th>( E_aR ) [K]</th>
</tr>
</thead>
<tbody>
<tr>
<td>( k_3 )</td>
<td>( 1.13 \times 10^{22} )</td>
<td>( 1.53 \times 10^4 )</td>
</tr>
<tr>
<td>( k_{14} )</td>
<td>( 8.13 \times 10^{16} )</td>
<td>( 1.28 \times 10^4 )</td>
</tr>
</tbody>
</table>

The dependence on temperature for the rate-determining step constants was determined at each pressure by applying the Arrhenius equation in the following way:

\[ \ln k = \frac{-E_a}{RT} - \ln k^0 + \ln k^0 \]

where \( T_0 \) is the mid-temperature for the temperature range considered, \( k^0 = A \times \exp(-E_a/R T_0) \), \( E_a \) is the activation energy, and \( A \) is the frequency factor.

The obtained results for the frequency factor and activation energy for each constant are reported in Table 7. The confidence interval limits of the frequency factor and activation energy are not reported because in many cases this confidence interval was greater than the value itself. This was obtained due to the poor fitting of the rate constants with respect to Arrhenius equation.
However, this lack of fitness is attributed to the experimental error obtained during the collection of data, and also to the shortage of experiments at different temperatures. Only three temperatures were analyzed, which are not sufficient to have conclusive results; therefore, more experiments at different temperatures within the studied interval should be performed.

The same kinetic mechanism was employed to determine the kinetic parameters for the hydrogenation of cyclohexene using the catalyst RhCl(TAN15DPPA)3 with 30% of oxidized phosphines. The temperature range studied was 70–120 °C at 172 bar. The substrate to catalyst molar ratio was 400 at 70 °C, and 700 at 95 and 120 °C. The initial mole fraction of reactants was kept constant in all the experiments at different temperatures.

In this case, the formation of monohydride catalytic species is taken into consideration even though no isomerization products were detected. This was done in order to keep the same kinetic mechanism. The double bond in cyclohexene may be shifting around, but it does not produce a distinct new isomer.

If reaction 3 is considered the rate controlling for hydrogenation, then the following kinetic equations are obtained:

$$
\frac{dC_A}{dt} = \frac{k_3K_1K_2C_A C_B C_{cat}}{1 + (1 + K_2C_A)K_1C_B + K_5^{1/2}C_B^{1/2}}
$$

$$
\frac{dC_{cat}}{dt} = \frac{k_{14}C_A C_{cat}}{1 + (1 + K_2C_A)K_1C_B + K_5^{1/2}C_B^{1/2}} - k_{14}C_{cat}
$$

where

$$
A = \text{cyclohexene}
$$

$$
B = \text{hydrogen}
$$

$$
P = \text{cyclohexane}
$$

Table 8 presents the values of the estimated parameters, and Figure 15 shows the comparison between the
calculated and the experimental results. Once more, the calculated values fit very well the experimental data.

Again, the dependence on temperature of the rate-determining step constant was determined using the Arrhenius equation. The results are presented in Table 9.

One more time, lack of fitness of rate-determining step constant with respect to the Arrhenius equation was obtained for the same reasons explained above.

Conclusions

A mechanism cycle was proposed for the hydrogenation and isomerization of 1-octene, and for the hydrogenation of cyclohexene. Statistical methods were applied to discriminate among rival models. It was proposed that two hydride catalytic species are formed during the reaction cycle: dihydride species for promoting hydrogenation, and monohydride species for yielding isomerization. Also, deactivation of the catalyst was attributed to the formation of unsaturated olefin complex on the catalyst. It was found that the rate-determining step for hydrogenation was the migratory insertion of hydrogen into the olefin to produce an alkylrhodium hydride complex. Moreover, the rate-determining step for isomerization was considered to be the coordination of the olefin in the monohydride catalytic species. The final kinetic model was found to be consistent with the general trends of the experimental data for both the hydrogenation and isomerization of 1-octene and cyclohexene.

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