**Hydrogenation of the Liquid Organic Hydrogen Carrier Compound Dibenzyltoluene – Reaction Pathway Determination by 1H-NMR Spectroscopy**

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<th>Journal:</th>
<th>Reaction Chemistry &amp; Engineering</th>
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<tr>
<td>Manuscript ID</td>
<td>RE-ART-11-2015-000080.R2</td>
</tr>
<tr>
<td>Article Type:</td>
<td>Paper</td>
</tr>
<tr>
<td>Date Submitted by the Author:</td>
<td>03-Feb-2016</td>
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</table>
| Complete List of Authors: | Do, Giang; Chemical Reaction Engineering, Department of Chemical and Bioengineering  
Preuster, Patrick; Chemical Reaction Engineering, Department of Chemical and Bioengineering  
Aslam, Rabya; Friedrich-Alexander-Universität Erlangen-Nürnberg, Chair for Separation Science and Technology  
Bösmann, Andreas; Chemical Reaction Engineering, Department of Chemical and Bioengineering, University Erlangen-Nuremberg  
Müller, Karsten; Friedrich-Alexander-Universität Erlangen-Nürnberg, Chair for Separation Science and Technology  
Arlt, Wolfgang; Friedrich-Alexander-Universität Erlangen-Nürnberg, Chair for Separation Science and Technology  
Wasserscheid, Peter; Chemical Reaction Engineering, Department of Chemical and Bioengineering |
DBT-hydrogenation pathway revealed by $^1$H-NMR

Chemical shift distributions (CS) for different samples:
- CS[7.0]
- CS[4.0]
- CS[2.0-2.5]
- CS[0.8-1.8]

Graphical representation of reaction products and their relative concentrations as a function of chemical shift ratio.

Legend:
- $T_{Reaction} = 120^\circ C$ - MSS
- $T_{Reaction} = 140^\circ C$ - Simult.
- $T_{Reaction} = 160^\circ C$ - Simult.
- $T_{Reaction} = 180^\circ C$ - Simult.
- $T_{Reaction} = 200^\circ C$ - Simult.
Hydrogenation of the Liquid Organic Hydrogen Carrier Compound Dibenzyltoluene – Reaction Pathway Determination by \(^1\)H-NMR Spectroscopy

G. Do\(^a\), P. Preuster\(^a\), R. Aslam\(^a\), A. Bösmann\(^a\), K. Müller\(^b\), W. Airt\(^b\) and P. Wasserscheid\(^{a,c,f}\)

The catalytic hydrogenation of the LOHC compound dibenzyltoluene (H0-DBT) was investigated by \(^1\)H-NMR spectroscopy in order to elucidate the reaction pathway of its charging process with hydrogen in the context of future hydrogen storage applications. Five different reaction pathways during H0-DBT hydrogenation were considered including middle-ring preference (middle-side, MMS), side-middle-side order of hydrogenation (SMS), side-ring preference (SSM), simultaneous hydrogenation of all three rings without intermediate formation and statistical hydrogenation without any ring preference. Detailed analysis of the \(^1\)H-NMR spectra of the H0-DBT hydrogenation over time revealed that the reaction proceeds with a very high preference in the SSM order at temperatures between 120 °C and 200 °C and 50 bar in presence of a Ru/Al\(_2\)O\(_3\)-catalyst. HPLC-analysis supported this interpretation by confirming an accumulation of H12-DBT-species prior to full hydrogenation to H18-DBT with middle ring hydrogenation as the final step.

Introduction

Hydrogen can be produced from renewable electricity via electrolysis and its energetic use (e.g. in a fuel cell or a hydrogen combustion engine) forms water as the only product. This enables a carbon- and therefore CO\(_2\)-free storage cycle for renewable energy equivalents with enormous storage capacity. However, while gravimetric energy storage density of hydrogen is excellent (120 MJ/kg or 33.3 kWh/kg), its volumetric energy density is a critical aspect. At ambient conditions it is only 3 Wh/liter. Even in compressed form (CGH\(_2\), 700 bar H\(_2\)) and in liquefied cryogenic form (LH\(_2\), liquid H\(_2\) at -253 °C) it is only 1.3 kWh/liter and 2.4 kWh/liter, respectively. In addition, hydrogen compression and cooling are energy intensive steps that require special infrastructure, such as compressor stations or cryogenic tank systems.

“Liquid Organic Hydrogen Carriers” (LOHCs) systems represent a relatively recent and very promising concept to establish a widespread hydrogen storage and transport system using the existing infrastructure for liquid fuels. LOHC systems typically comprise a pair of organic molecules, a hydrogen-lean and a hydrogen-rich compound. Both are liquids and their physico-chemical properties highly resemble conventional fuels. Other important selection criteria for suitable LOHC systems are derived from economic, safety, environmental and health impact assessments.

During the hydrogen storage process, the hydrogen-lean LOHC compound is transformed into the hydrogen-rich compound by a catalytic hydrogenation reaction. The hydrogenated compound of the LOHC system can be stored and transported in large quantities with no energy losses over time for a later release of hydrogen and energy on demand, i.e. wherever and whenever energy or hydrogen is most valuable. It is a key aspect of the LOHC technology that the LOHC material is used in many repeated hydrogenation/dehydrogenation cycles and acts like a liquid container for the stored hydrogen. No gaseous reactants other than H\(_2\) are needed in the hydrogen storage process and no coupling products besides the liquid carrier are released during hydrogen generation from LOHC systems. This distinguishes the LOHC technology in a fundamental manner from other “power to X” or “power to gas” technologies using nitrogen or CO\(_2\) as reactants to form, e.g. ammonia, methane or methanol, as storage compounds.

Among the known LOHC systems, the system dibenzyltoluene (H0-DBT)/perhydro-dibenzyltoluene (H18-DBT) is of particular interest. Dibenzyltoluene is a commercial heat transfer oil (marketed e.g. under the tradename Marlotherm SH©) with known and very favorable toxicology (non-toxic, non-mutagenic, non-carcinogenic, not labelled as hazardous good in transportation), high thermal stability and excellent physico-chemical properties for many LOHC application scenarios. For example the isomeric mixture of dibenzyltoluenes comes with a melting point of -34 °C and a boiling point of 390 °C. During hydrogenation, H0-DBT binds 6.2 mass% H\(_2\) corresponding to an energy content of 2.05 kWh/kg.

In this contribution we report on a series of investigations aiming for a more detailed mechanistic understanding of the heterogeneously catalyzed hydrogenation of H0-DBT to H18-DBT. This reaction represents one key step in the practical use of the respective LOHC system and determines the dimensions of the required hydrogenation reactor (including investment costs) but also reliability and robustness of the hydrogen loading process onto the LOHC material. Therefore, it is of significant interest to optimize catalyst materials, reaction conditions and reaction devices for this reaction. An important aspect for these optimization efforts is to know more about the reaction pathway of the catalytic hydrogenation. As H0-
DBT is characterized by three aromatic groups in the molecules, the order of hydrogenation of these three is of interest.

Studies in the field of hydrogenation of poly aromatic systems have shown different selectivities\textsuperscript{12-15} towards the partially hydrogenated intermediates depending on the choice of catalyst system and reaction conditions. Crompton showed that the primary intermediates of o- and m-terphenyl hydrogenation using PtO\textsubscript{2} are the middle ring saturated species\textsuperscript{16}. Similar observations were made by Scola using copper chromite catalyzed reduction of o-terphenyl\textsuperscript{17} which led to a 10:1 ratio of center ring to outer ring reduction. However, hydrogenating m-terphenyl led to a ratio of 1.6:1. The use of a nickel catalyst showed a definite preference for outer-ring reduction. The known analytic methods for poly aromatic systems are HPLC,\textsuperscript{18, 19} GC-analysis\textsuperscript{17, 20, 21} or detection via ultraviolet spectroscopy\textsuperscript{22} involving a separation step of the various species. In the present paper we elaborate the hydrogenation and dehydrogenation sequences for the H0-DBT/H18-DBT system by using \textsuperscript{1}H-NMR spectroscopy\textsuperscript{23-25} in combination with HPLC and GC-MS analyses as well as suitable kinetic models. For H0-DBT hydrogenation this combination of analytic methods allows for an unambiguous determination of the dominating reaction pathway.

**Experimental**

**Materials:** H0-DBT was purchased from Sasol. The applied commercial Ru/Al\textsubscript{2}O\textsubscript{3} catalyst was supplied by Hydrogenious Technologies (www.hydrogenious.net).

**Hydrogenation:** The hydrogenation experiments were carried out in a 300 mL stainless-steel batch auto-clave (Parr Instrument Company) with a four-blade gas-entrainment stirrer (\( n = 1200 \) rpm). First, 150 g H0-DBT was loaded into the vessel together with the catalyst at a molar ratio of 400 : 1 (H0-DBT/Ru). After purging the gas volume three times with Ar 4.6 (argon with a purity of 99.999%), the vessel was heated with an electrical heating jacket to the reaction temperature. The pressure was then adjusted and held constant at 50 bar during the experiment. In defined time intervals liquid samples were taken to determine the progress of hydrogenation.

**NMR-Analysis:** Nuclear magnetic Resonance (NMR) was applied in the determination of the reaction mixture during the hydrogenation progress. For this purpose an ECX 400 from JEOL was applied. 0.1 mL of liquid sample was diluted in 1 mL of dichloromethane-d2 for sample preparation.

**HPLC analysis:** Reverse phase HPLC analysis was performed using a Merc-Itachi Elite Lachrom HPLC-system equipped with an auto sampler, degasifier, mixer and L-7100 HPLC pumps, UV-vis and RI detectors. Chromatographic separation of the fractions based on the degree of hydrogenation, i.e. in fractions of H0-DBT, H6-DBT, H12-DBT, and H18-DBT was achieved on a Phenomenex phenyl-hexyl silica column (250 mm x 4.6 mm I.D, 15 \( \mu \)m, pore size 100 Å). Acetone and water (95:5, v/v) were used as eluents in isocratic mode with flow rate of 0.17 ml/min. The separated fractions were analyzed with GC MS to determine their composition. Samples from the reactor were taken and analyzed with HPLC to determine the composition change during the hydrogenation reaction of H0-DBT to H6-, H12-, and H18-DBT as a function of time.

**Results and discussion**

**Reaction pathway.** Assuming that species with partially hydrogenated rings and regioisomers can be neglected, up to 6 DBT-species may occur during the hydrogenation progress from H0-DBT to H18-DBT. Close examination reveals that H18-DBT can be reached via five different reaction pathways (see Figure 1). Note that all depicted Hx-DBT structures stand for the different possible regioisomers (different relative positions of the benzyl groups vs. the methyl substituent at the central aromatic ring). For clarity only one regioisomer per DBT-species is shown in Figure 1. The first three reaction pathways (MSS, SMS and SSM) exhibit different hydrogenation preferences of the three aromatic rings of H0-DBT. While the simultaneous reaction path has a lack of intermediates, the statistical path exhibits no ring preference in the hydrogenation process.
Estimation of chemical shifts. In $^1$H-NMR, hydrogen atoms surrounded by different chemical functionalities show different shifts. According to the various Hx-DBT species formed during the hydrogenation process of H0-DBT (see Figure 1), the corresponding chemical shifts move from the aromatic regime over the mixed aromatic/aliphatic regime to the purely aliphatic regime. Using the ChemBioDraw software, $^1$H-NMR spectra of all structural isomers of each Hx-DBT species were predicted in order to define the respective ranges for signal integration. Four different ranges representing different types of hydrogen atoms can be distinguished for the H0-DBT hydrogenation process. As an example, a single-side-ring hydrogenated H6-DBT molecule is shown together with the corresponding NMR-spectrum and chemical shifts in Figure 2. As an example, the predicted $^1$H-NMR spectra of all H0-DBT isomers are shown in Figure 2 together with the respective measured H0-DBT sample. For all regioisomers, the hydrogen atoms with different chemical shifts can directly be associated to the assigned chemical shift ranges shown in Table 1.

Figure 2: $^1$H NMR spectra and respective chemical shift ranges of all H0-DBT structural isomers compared with measured educt sample H0-DBT: a) Measured educt sample of H0-DBT; b) 2,3-dibenzyltoluene; c) 3,4-dibenzyltoluene; d) 2,6-dibenzyltoluene; e) 2,4-dibenzyltoluene; f) 3,5-dibenzyltoluene; g) 2,5-dibenzyltoluene.

Table 1: Chemical shifts of selected hydrogen atoms in $^1$H-NMR spectroscopy – R represents the rest of the Hx-DBT-species; hydrogen atoms with chemical shifts outside the indicated range are not shown explicitly.

<table>
<thead>
<tr>
<th>Chemical shift</th>
<th>Corresponding structural environment</th>
</tr>
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<tr>
<td>ca 7.0 ppm</td>
<td>(CS[7.0])</td>
</tr>
<tr>
<td>ca 4.0 ppm</td>
<td>(CS[3.9-4.1])</td>
</tr>
<tr>
<td>2.0-2.5 ppm</td>
<td>(CS[2.0-2.5])</td>
</tr>
<tr>
<td>0.8-1.8 ppm</td>
<td>(CS[0.8-1.8])</td>
</tr>
</tbody>
</table>

Integration of the four shift ranges and calculation of the fractions according to Equations 1 to 4 has been carried out to determine the fractions of the different types of hydrogen atoms in the respective Hx-DBT samples. According to Equation 5, the sum of all fractions for the different types of hydrogen atoms is 1. Thus, if three fractions are known, the forth can be calculated according to Equ. 5. For this study, the fractions of %CS[7.0], %CS[4.0] and %CS[0.8-1.8] have been determined from the NMR-spectra and were used for data evaluation and interpretation.

\[
\frac{\%CS[7.0]}{100} = \frac{j_{\text{CS}} f_{\text{CS}}}{j_{R} f_{R}}
\]

(1)

\[
\frac{\%CS[4.0]}{100} = \frac{j_{\text{CS}} f_{\text{CS}}}{j_{R} f_{R}}
\]

(2)

\[
\frac{\%CS[2.0-2.5]}{100} = \frac{j_{\text{CS}} f_{\text{CS}}}{j_{R} f_{R}}
\]

(3)

\[
\frac{\%CS[0.8-1.8]}{100} = \frac{j_{\text{CS}} f_{\text{CS}}}{j_{R} f_{R}}
\]

(4)

\[
\%CS[7.0] + \%CS[4.0] + \%CS[2.0-2.5] + \%CS[0.8-1.8] = 1
\]

(5)
The fractions of the chosen chemical shift integrals were calculated for all Hx-DBT-species and are shown in Table 2. Pairing two of the three chemical shift fractions leads to three different plots, showing the characteristic fractions of $^1$H-NMR shifts for the respective reaction paths from Figure 1. These predictions are displayed in Figure 3 to 5.

Some aspects should be highlighted here to better understand Figures 3 to 5:

- **Simultaneous H0-DBT hydrogenation exhibits only two species according to Figure 1.** The progress for statistical hydrogenation has been calculated by taking into account a one third chance of middle ring and a two third chance of side ring hydrogenation according to the share of hydrogen to be transferred to these two different positions.

- **Hydrogen atoms at the methylene bridges without any aromatic environment exhibit no signal at the CS[4.0] regime.** Thus, hydrogenation preference for the middle ring of DBT shows a steeper decrease of CS[4.0]-signals compared to any pathway preferring side-ring hydrogenation in the plots of Figure 3 and Figure 4.

- **The hydrogenation with side-ring preference shows the flattest slope between H0- and H12-species in the plot in Figure 5 as the number of aromatic hydrogen atoms being converted to aliphatic hydrogens through side ring saturation is higher compared to the middle ring hydrogenation.**

### Table 2: Fractions of chemical shift integrals

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>H0-DBT</th>
<th>H6-DBT</th>
<th>H6-DBT</th>
<th>H12-DBT</th>
<th>H12-DBT</th>
<th>H18-DBT</th>
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<tr>
<td>Degree of Hydrogenation</td>
<td>0%</td>
<td>33.3%</td>
<td>33.3%</td>
<td>66.7%</td>
<td>66.7%</td>
<td>100%</td>
</tr>
<tr>
<td>Molecular structure</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>%-CS[7.0]</td>
<td>0.65</td>
<td>0.31</td>
<td>0.38</td>
<td>0.16</td>
<td>0.09</td>
<td>0.00</td>
</tr>
<tr>
<td>%-CS[4.0]</td>
<td>0.20</td>
<td>0.08</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
</tr>
<tr>
<td>%-CS[0.8-1.8]</td>
<td>0.00</td>
<td>0.42</td>
<td>0.46</td>
<td>0.78</td>
<td>0.69</td>
<td>1.00</td>
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![Figure 3](image_url)  
**Figure 3:** Calculated fractions of chemical shift integrals of hydrogen atoms of the methylene bridge connecting two aromatic rings (chemical shift at 4 ppm) and aliphatic hydrogen atoms (chemical shift below 1.8 ppm) for all Hx-DBT species. SSM: Side-ring preference; SMS: side-ring, middle-ring, side-ring; MSS: Middle-ring preference; simultaneous: hydrogenation of all three rings at the same time; statistical: hydrogenation without preferences.

![Figure 4](image_url)  
**Figure 4:** Calculated fractions of chemical shift integrals of hydrogen atoms of the methylene bridge connecting two aromatic rings (chemical shift at 4 ppm) and aromatic hydrogen atoms (chemical shift at 7 ppm) for all Hx-DBT species. SSM: side-ring preference; SMS: side-ring, middle-ring, side-ring; MSS: middle-ring preference; simultaneous: hydrogenation of all three rings without intermediates; statistical: hydrogenation without preferences.
Figure 5: Calculated fractions of chemical shift integrals of aliphatic hydrogen atoms (chemical shift below 1.8 ppm) and aromatic hydrogen atoms (chemical shift at 7 ppm) for all Hx-DBT species. SSM: side-ring preference; SMS: side-ring, middle-ring, side-ring; MSS: middle ring preference; simultaneous: hydrogenation of all three rings at the same time; statistical: hydrogenation without preferences.

Evaluation of the potential reaction path “Simultaneous”. For an estimated “simultaneous” reaction pathway, the reaction mixture consists of a mixture of H0-DBT and H18-DBT as no intermediate hydrogenation products (e.g. H6-DBT or H12-DBT) are formed in this mechanism. To check this option, artificial reaction mixtures were prepared by mixing eleven different fractions of H0-DBT and H18-DBT and these mixtures were measured by $^1$H-NMR. Ratios of chemical shift integrals are plotted in Figure 6 to Figure 8 for these mixtures to compare the fractions of chemical shifts obtained in catalytic hydrogenation experiments with the expected trend for such a simultaneous hydrogenation.

Reaction path of the H0-DBT-hydrogenation. By evaluating the fractions of chemical shift integrals of samples taken during a catalytic H0-DBT hydrogenation experiment, the appearance of these fractions in the NMR spectra are obviously not influenced by the reaction temperature, as shown in Figure 9 to 11. This reveals that in the investigated temperature range between 120 and 200°C the applied Ru on Al$_2$O$_3$ catalyst hydrogenates H0-DBT in the same order of aromatic rings. Furthermore, based on the result in Figure 9 the reaction paths following the MSS hydrogenation order as well as the Simultaneous and Statistical pathway can be excluded. Simply, the calculated ratios of CS[4.0] vs. CS[0.8-1.8] as shown in Figure 9 are too far off the experimental values measured for the real hydrogenation process. Taking additionally Figure 10 into account, the reaction path SMS becomes very unlikely while the experimental data match particularly well the reaction path SSM.
The same conclusion of a strongly preferred SSM reaction path is also confirmed by Figure 11 where the share of bridging CH$_2$ neighboring by fully hydrogenated cyclohexyl groups is plotted against the share of aromatic protons.

Concerning the revealed SSM pathway, there exist three different options for the pathway to proceed as shown in Figure 12. All steps may happen in sequence resulting in a three step pathway or two of the respective steps may happen simultaneously. [SS]M refers to simultaneous hydrogenation of both side-rings as the first step and the middle-ring hydrogenation as the second step, whereas [S]M represents a single side-ring hydrogenation as the first step and simultaneous side-ring and middle-ring hydrogenation as the second step. Thus, pathway [SS]M lacks of H6-DBT$_{XOO}$ and [S]M lacks of H12-DBT$_{XXO}$. NMR results reveal that H12-DBT$_{XOO}$ must exist during the hydrogenation process, due to the fact that in all chemical shift integral fractions of Figure 9 to 13, H12$_{XOO}$ is confirmed by experimental data. Lacking of H12$_{XOO}$ would lead to a linear connection line between H6$_{XOO}$ and H18$_{XXO}$. Hence S(SM) can be excluded.

On the contrary, NMR results are not conclusive to exclude either [SS]M or SSM, because in both cases the hydrogenation progress goes from H0$_{XOO}$ to H12$_{XOO}$ via H6$_{XOO}$. Since mixtures of H0$_{XOO}$ and H12$_{XOO}$ can also represent H6$_{XOO}$, NMR experimental data are not sufficient here for the final discrimination. Therefore, HPLC investigations of Hx-DBT reaction mixtures were also performed. According to the mass-percentage of all Hx-DBT species plotted over hydrogenation time shown in Figure 13, the three-step SSM reaction is predominant due to the presence of H6- and H12-species. In addition, the main fraction of H18-species emerges when H12 accounts for ~90wt-% indicating that middle ring hydrogenation only starts when most of the side rings are saturated. Accordingly, the combination of NMR-spectroscopy and HPLC reveals unambiguously a SSM hydrogenation sequence with three independent hydrogenation steps.
Conclusions

In this contribution we report a detailed analysis of the catalytic hydrogenation of dibenzyltoluene on a Ru on Al₂O₃ catalyst using ¹H-NMR spectroscopy and HPLC. The reaction is of high relevance in the context of using the pair dibenzyltoluene (H₀-DBT)/perhydro dibenzyltoluene (H₁₈-DBT) as liquid organic hydrogen carrier (LOHC) system. It has been found that in the studied parameter range (T = 120-200 °C, 50 bar hydrogen) the reaction proceeds with a very high preference in the way that first the two side phenyl rings and last the middle phenyl ring are hydrogenated. This new mechanistic insight may help to interpret kinetic results from the H₀-DBT hydrogenation and to understand macrokinetic effects with the applied porous catalyst system. Moreover, the methodology presented is suitable to compare different reaction conditions and catalyst materials with regard to the sequence of the hydrogenation pathway. Such insight may help to identify mechanistically different H₀-DBT hydrogenation catalysts in the course of further optimization studies of the H₀-DBT catalyst. It is an interesting option, for example, to combine within the hydrogenation catalyst bed layers of different catalysts. Based on the here presented results, Ru on Al₂O₃ appears very suitable for fast side phenyl ring hydrogenation. For finishing the final middle ring hydrogenation a make-up catalyst layer with a different catalyst material could be more suitable, as it has been found, that middle ring hydrogenation with Ru on Al₂O₃ is a slow step that only sets in when almost every LOHC has been converted to the H₁₂-DBT state. Most remarkably, no side products outside the intended reaction pathway from H₀-DBT towards H₁₈-DBT have been detected at least by NMR. This confirms that hydrogen storage in the LOHC system H₀-DBT / H₁₈-DBT is indeed a very selective process making this system highly suitable for repeated hydrogen storage in reversible hydrogenation/dehydrogenation cycles.

Acknowledgements

The authors acknowledge financial support by the application center VerTec in Fürth, by the Bavarian Hydrogen Center, by the Energie Campus Nürnberg and by the Erlangen Excellence Cluster "Engineering of Advanced Materials”.

Notes and references


