Separation Enhanced Dimethyl Ether Synthesis

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A novel sorption enhanced dimethyl ether synthesis (SEDMES) process is presented using a solid adsorbent to remove produced water in situ. SEDMES experiments from feed mixtures of \( \text{H}_2 \), \( \text{CO} \), and \( \text{CO}_2 \) have shown an increased yield of DME, an improved selectivity to DME over methanol, and a strongly reduced \( \text{CO}_2 \) content in the product. Consequently, SEDMES will reduce the downstream separation effort and minimise the recycle streams. Within the European Horizon 2020 project FLEDGED, synthesis gas from biomass gasification will be used as feedstock for the separation enhanced DME-synthesis.

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**Introduction**

Dimethyl ether (DME) is a promising alternative fuel among the various ultra clean, renewable, and low-carbon fuels under consideration worldwide [1]. In the DME production from syngas, the following equilibrium reactions are involved:

- Methanol synthesis
  \[ \text{CO}_2 + 3\text{H}_2 \rightleftharpoons \text{CH}_3\text{OH} + \text{H}_2\text{O} \quad (1) \]
- Water-gas shift (WGS)
  \[ \text{CO} + \text{H}_2\text{O} \rightleftharpoons \text{H}_2 + \text{CO}_2 \quad (2) \]
- Methanol dehydration
  \[ 2\text{CH}_3\text{OH} \rightleftharpoons \text{CH}_3\text{OCH}_3 + \text{H}_2\text{O} \quad (3) \]

Indirect DME production comprises the production of intermediate methanol (1), and methanol dehydration (3) in separate reactors. Both reactions are thermodynamically limited which leads to limited DME yield and extensive separations and recycles. The direct DME synthesis (Figure 1b) proceeds in a single reactor via intermediate methanol (1) as well, yet offers a reduction in process steps and increased overall DME yield. In terms of efficiency, direct DME synthesis process outperforms the indirect synthesis, yet the need for separation and recycling remains. In the direct DME synthesis, the \( \text{O} \)-surplus of the feed ends up in \( \text{CO}_2 \), which means that about equal molar amounts of DME and \( \text{CO}_2 \) are produced. Since the reaction is equilibrium limited, downstream separation produces recycle streams of syngas (\( \text{CO} \) and \( \text{H}_2 \)), \( \text{CO}_2 \), and methanol. Syngas and methanol are recycled back to the DME synthesis reactor, while the \( \text{CO}_2 \) recycle is used in synthesis gas generation via dry or tri-reforming.

A novel process route exists, called sorption enhanced DME synthesis (SEDMES, Figure 1c). It is based on the use of a solid adsorbent for the in situ removal of water. According to Le Chatelier’s principle, the removal of one of the products will shift the equilibrium-limited conversion to the product side. The process has been analysed theoretically [2], indicating that in situ water adsorption leads to an increased yield and selectivity to DME. In this contribution, results are presented of an investigation into the sorption-enhanced production of DME.

**Approach**

A combined approach was taken, comprising thermodynamic modelling and transient experiments. Thermodynamic calculations were performed on different feed compositions with different \( \text{H}_2/\text{CO}/\text{CO}_2 \) ratios (see Table 1) using HSC Chemistry [3]. All compositions had an \( M \) module of 2, the stoichiometric molar ratio for DME, defined as \( M = ([\text{H}_2]-[\text{CO}_2])/([\text{CO}]+[\text{CO}_2]) \). The calculations were performed taking into account equilibrium reactions (1-3) from the feed mixtures at 25 bar(a) pressure and as a function of temperature. In addition, the impact of steam separation enhanced reaction was addressed by calculating the thermodynamic equilibrium as a function of...
steam content at a total pressure of 25 bar(a) and a temperature of 150-350 °C.

Experimentally, a combination of a commercial copper/zinc oxide/alumina catalyst and a commercial zeolite 3A steam adsorbent was used for the demonstration of direct DME synthesis from H₂/CO/CO₂ mixtures. Particles were mixed as sieve fractions. The experimental runs were conducted on a high-pressure reactor setup, allowing 50 g of sample, consisting of 10 g of catalyst and 40 g of sorbent, to be tested in a single reactor of 15.75 mm internal diameter. The catalyst to adsorbent ratio was arbitrarily chosen and not further optimised in this work. Adsorption was done under different feed gas compositions, shown in Table 1. The N₂ balance was used in order to keep the overall pressure stable, considering the shrinking overall reaction. Regeneration was done by switching to dry nitrogen, depressurisation to 1.7 bar(a), and heating to 400 °C. Gas analysis was done by i) MS measuring H₂ (m/z=2), H₂O (m/z=18), CO/N₂ (m/z=28), CH₃OH (m/z=31), CO₂ (m/z=44) and DME (m/z=45), and ii) NDIR measuring CO and CO₂ in the dry sample flow.

Fig. 2: Thermodynamic equilibrium versus temperature for the compositions in Table 1, 25 bar(a)

Fig. 3: Thermodynamic equilibrium versus water slip level for the compositions in Table 1, 25 bar(a), 275 °C

Results
Figure 2 shows the thermodynamic results on the formation of DME as a function of temperature for the gas compositions given in Table 1, without in situ water adsorption. DME formation is dependent on the different H₂/CO/CO₂ ratios as well as temperature. At decreasing oxygen content in the gas mixture (order of composition 1, 3, and 2) the conversion to DME is higher, indicating that the presence of excess oxygen is hampering the formation of DME. Oxygen removal can be realised by removing the reaction product steam and thus promote the conversion to DME. This is essentially the removal of steam from the reactions (1-3). Figure 3 shows the results of the thermodynamic calculations of the formation of DME as a function of temperature. The partial pressure of each gas in the composition was chosen to achieve thermodynamic equilibrium.
of steam content at 25 bar(a) for the three gas compositions given in Table 1. Because \( M = 2 \) for all three cases, the graphs appear very similar and clearly show that the removal of steam results in higher conversions towards DME for all gas compositions. The carbon conversion to DME is very high, especially at lower steam slip levels. The concentrations of methanol and CO\(_2\) reduce significantly, suggesting that the conversion of oxygen rich feedstock CO\(_2\) into DME can be realised when removing steam.

![Fig. 4: Breakthrough experiment with Composition 3, 25 bar(a), 275 °C](image)

![Fig. 5: Conventional versus sorption enhanced direct DME synthesis, 25 bar(a), 275 °C](image)

Experimental proof-of-concept of SEDMES is indicated in Figure 4 by breakthrough measurement of Composition 3. Before the breakthrough of steam, which starts at about 16 minutes on stream, the main carbon-containing products are CO and DME. A small amount of methanol is being produced, and small amounts of H\(_2\)O and CO\(_2\) slip through the not yet optimised bed. After 16 minutes on stream, H\(_2\)O and CO\(_2\) break through together and the DME concentration drops. Similar results were obtained with the other feed gas compositions, and the observed pre-breakthrough sorption enhancement for each of the settings is shown in Figure 5. The amount of feed C ending up in DME is enhanced from 4-50% for the direct DME synthesis to around 65% for SEDMES. Additionally, the amount ending up in CO\(_2\) is strongly reduced from 20-80% to below 2%. Moreover, the product distribution depends much less on the feed CO/CO\(_2\) composition (at the same value of the \( M \) module). When comparing the obtained pre-breakthrough concentrations (Figure 5) to the thermodynamic equilibrium (Figure 3), it shows that the experimental product spectrum is according to thermodynamic equilibrium. It is anticipated, therefore, that removal of balance N\(_2\) will further enhance the conversion to DME. These observations for the sorption enhanced mode have several important implications for DME synthesis. The feed to the SEDMES reactor can contain CO\(_2\) as well as CO. Secondly, the CO\(_2\) content of the reactor effluent is largely suppressed, simplifying the downstream separation section and largely eliminating the CO\(_2\) recycle. These findings have led to a recent patent application on SEDMES [4].

**Summary**

A novel sorption enhanced dimethyl ether synthesis (SEDMES) process is presented using a solid adsorbent to remove produced water in situ. SEDMES experiments from feed mixtures of H\(_2\), CO, and CO\(_2\) have shown an increased yield of DME, an improved selectivity to DME over methanol, and a strongly reduced CO\(_2\) content in the product. Consequently, SEDMES will reduce the downstream separation effort and minimise the recycle streams.

**Outlook**

Fledged is a recently started EU H2020 project which aims at developing a flexible process for the production of DME from biomass gasification with sorption enhanced processes. In Fledged, the SEDMES process is being developed further, targeting higher conversions to DME. Specific attention is paid to elucidation of the reaction mechanism. The objective of the project is an experimental validation of a full cyclic SEDMES process for 3 kg/h DME production at ECN laboratories.

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**References**


