Sorption enhanced dimethyl ether synthesis under industrially relevant conditions: experimental validation of pressure swing regeneration

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Dimethyl ether (DME) is one of the most attractive alternative fuel solutions under consideration worldwide. However, its production from CO₂-rich feedstock or CO₂ directly is limited via conventional processes and therefore considered unattractive. For CO₂ utilisation, the production and efficient handling of steam remains a major bottleneck. Sorption enhanced DME synthesis (SEDMES), which combines heterogeneous catalysis with in situ water adsorption, is a promising process intensification strategy for the direct production of DME from CO₂. In this work, SEDMES is demonstrated experimentally on a bench-scale reactor with pressure swing regeneration under industrially relevant conditions. Pressure swing regeneration, rather than the time and energy intensive temperature swing regeneration, shows high performance with over 80% single-pass carbon selectivity to DME. This already allows for a factor four increase in productivity, with further optimisation still possible. With the proposed Sips working isotherm for the water adsorbent, and the methanol synthesis and dehydration kinetics, the validated dynamic cycle model adequately describes the SEDMES bench-scale data. Applying shorter cycle times, made possible by pressure swing regeneration, allows optimisation of the DME productivity while maintaining the high single-pass yield typical for SEDMES. The experimental confirmation shown in this paper unlocks the full potential of the high efficiency carbon and hydrogen utilisation by SEDMES technology.

1. Introduction

Dimethyl ether (DME) is the simplest ether compound, with a chemical formula of CH₃OCH₃. DME is gaseous at ambient conditions, which is easy to liquefy and transport. It is safely stored and handled and does not form explosive peroxides in contrast to several other ethers. Its chemical and physical properties as well as its combustion characteristics make that DME can be used as fuel in domestic applications replacing LPG, in compression ignition engines (100% DME) and spark ignition engines (30% DME/70% LPG), and in power generation. Consequently, DME is one of the most promising alternative fuel solutions among the various ultra clean, renewable, and low-carbon fuels under consideration worldwide.¹⁻³

Conventionally, DME is produced from synthesis gas with methanol as an intermediate chemical. The following equilibrium reactions are involved:

\[ \text{Methanol synthesis} \quad \text{CO} + 3\text{H}_2 \rightleftharpoons \text{CH}_3\text{OH} + \text{H}_2\text{O} \quad (1) \]

\[ \text{Methanol synthesis} \quad \text{CO} + 2\text{H}_2 \rightleftharpoons \text{CH}_3\text{OH} \quad (2) \]

\[ \text{Water–gas shift} \quad \text{CO} + \text{H}_2\text{O} \rightleftharpoons \text{H}_2 + \text{CO}_2 \quad (3) \]

\[ \text{Methanol dehydration} \quad 2\text{CH}_3\text{OH} \rightleftharpoons \text{CH}_3\text{OCH}_3 + \text{H}_2\text{O} \quad (4) \]

Indirect DME production comprises the production of intermediate methanol (1, 2), and methanol dehydration (4). The incomplete methanol and DME yields require extensive separation sections and recycles. The single-step direct DME synthesis proceeds via intermediate methanol as well, yet offers a reduction in process steps and increased overall conversion to DME.⁴⁻⁷ Although the direct DME synthesis process outperforms the indirect process in terms of efficiency, separation and recycling remain a requirement. In the direct DME synthesis, the O-surplus of the feed ends up in CO₂, resulting in equal molar amounts of DME and CO₂ produced. Since the reaction is equilibrium limited, the downstream separation section produces recycle streams of synthesis gas (CO + H₂), CO₂, and methanol. Synthesis gas and methanol can be recycled back to the DME synthesis reactor, while the CO₂ is at best
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2. Experimental

Materials

Experimental validation of sorption enhanced DME synthesis was performed using (a homogeneous mixture of) commercially available catalyst and adsorbent: CZA catalyst, γ-Al2O3 (assay >98%, Riogen NJ, USA), obtained as 3 mm pellets, and molecular sieve type 3A purchased as 1.6 mm pellets (UOP Molecular Sieves, Advanced Specialty Gas Equipment, USA).

Methods

The commercially obtained zeolite 3A adsorbent was tested at both atmospheric and high-pressure setups to determine its working capacity at relevant conditions. Fig. 1 represents the ‘Microflow 5’ test-rig for atmospheric experiments. A quartz reactor with an internal diameter of 10 mm was filled with 0.1–2 gram of material. During adsorption 100 mlN min\(^{-1}\) was fed to the reactor at 200–300 °C. The gas mixture contained 5–40 mol% H\(_2\)O, 5% CH\(_4\) as tracer and balance N\(_2\). Regeneration is performed by switching the gas flow to dry N\(_2\) and, in some cases, increasing the temperature to 350 °C for 5 minutes. Off-gas analysis was performed continuously by a Perkin Elmer Frontier FTIR with heated Pike 2.4 m gas cell.

In the next section, the used materials, the experimental procedures and model interpretation are reported. In the results and discussion section, firstly, the sorbent and catalyst performances are discussed separately for their input in the SEDMES model validation. This is followed by extensive SEDMES testing, model validation and prediction. Finally, the conclusions are summarized.
litres of sample, typically consisting of a 1:4 ratio (weight basis) catalyst to sorbent. The ratio between catalyst and sorbent was not further optimised in this work. Adsorption was performed with different (stoichiometric) feed gas compositions, using 68.6–72.7 vol% of hydrogen, 0–9.1 vol% of carbon monoxide, 17.1–23.6 vol% of carbon dioxide and

Fig. 1  Schematic presentation of the atmospheric and high pressure reactor units ‘Microflow 5’ and ‘Spider’.

Fig. 2  Schematic of the bench-scale reactor.
inert argon, nitrogen or methane, at 25 bar(a) and a temperature range of 250–300 °C. The inert balance was used in order to keep the overall pressure stable, considering the nett mole consumption by the reaction and the adsorption of water. Regeneration was done by depressurisation to 1–3 bar(a) for PSA regeneration, switching to dry, inert gas, and eventual heating to 400 °C for TSA regeneration. Finally, either the inert purge gas or the reactive feed gas is used for repressurisation. Transient gas analysis was performed by micro-GC (measuring methane, CO, CO₂, nitrogen, argon, methanol and DME) and mass spectrometry measuring hydrogen (m/z = 2), methane (m/z = 15), water (m/z = 18), carbon monoxide/nitrogen (m/z = 28), methanol (m/z = 31), carbon dioxide (m/z = 44) and DME (m/z = 45).

Data interpretation

To interpret the breakthrough experiments, a material balance for component i over the reactor column is made. Accumulation of component i between t = 0 and complete breakthrough (t = t_end) must be equal to the difference between the molar inflow and outflow rates.

\[
\frac{y_i(t_{\text{end}})pV_g}{RT} + q_i(t_{\text{end}})m_{\text{ads}} - \frac{y_i(0)pV_g}{RT} - q_i(0)m_{\text{ads}} = t_{\text{end}}F_{i,\text{in}} - \int_{t=0}^{t_{\text{end}}} (y_i F)_{\text{out}} \, dt
\]

The trapezoidal rule has been used to approximate the integral by a summation over discrete measurement data. The breakthrough of tracer (q_{tracer} = 0), prior to breakthrough of H₂O, is integrated to obtain V_g, the total interparticle and intraparticle gas volume. After breakthrough, the tracer signal is used to quantify the outlet flow rate prior to and during breakthrough and eqn (5) is then used to compute the water loading (q).

In order to facilitate data interpretation, several key metrics have been defined to be able to quantify the SEDMES performance. The most important metric, the carbon selectivity S(i), used here is defined as follows,

\[
S(i) = \frac{ny(C_{n}H_{m}O_{p})}{\sum n_{y}(i)}
\]

The carbon selectivities were calculated as molar concentration-based selectivities for each of the carbon containing species, y(i). For example, the selectivity towards DME can be calculated as

\[
S(\text{DME}) = \frac{2y(\text{DME})}{y(\text{CO}) + y(\text{CO}_2) + 2y(\text{DME}) + y(\text{MeOH}) + y(\text{CH}_4)}
\]

Time integration (in the interval t: 0 – t_{CO_2}, where t_{CO_2} is the (interpolated) point in time where the CO₂ outlet concentration reaches a level of 5 vol%) of the streams gives an overall yield and selectivity for the cyclic (steady state) performance of the SEDMES process.

Model

A one-dimensional pseudo-homogeneous dynamic reactor model was developed in Matlab, verified and validated. For the description of the fluid flow and mass transfer, the 1D non-steady differential mass and momentum balances are solved. The total mass, momentum, component and overall energy balances are given in Table 1. Reaction kinetics have been determined for the used catalyst materials by fitting the parameters in the models of Graaf et al. (1988) and Beričić et al. (1992) for the methanol synthesis and methanol dehydration respectively, shown in Table 2. The adsorption isotherm of the LTA zeolite adsorbent is determined under the high pressure and temperature working conditions of the SEDMES process. A Sips isotherm best describes the experimental data, in accordance with the available literature at lower temperature and pressure conditions. Full details of the different aspects of the model can be found in our previous work.

3. Results and discussion

To investigate the sorption enhanced DME synthesis process further, firstly the adsorption capacity at elevated temperature and pressure is looked into. This is followed by a performance study of the catalyst materials used in SEDMES. Proof-of-concept for sorption enhanced DME synthesis at bench-scale is demonstrated experimentally and

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**Table 1** Reactor model equations

| Equation                                      | Description                                                                 |
|----------------------------------------------|----------------------------------------------------------------------------|
| \( \frac{\partial \rho}{\partial t} = -\frac{\partial \rho v}{\partial z} - \frac{1-\varepsilon_b}{\varepsilon_b} a_p \sum M_i N_i \) | Overall mass balance                                                      |
| \( \frac{\partial \rho v}{\partial t} = -\frac{\partial \rho v^2}{\partial z} - \frac{\rho/\mu u}{\rho_p} \) | Momentum balance                                                          |
| \( \frac{\partial y_{oi}}{\partial t} = \frac{\partial y_{oi} v_{oi}}{\partial z} + \frac{\partial}{\partial z} \left( D_{oi} \frac{\partial y_{oi}}{\partial z} \right) - \frac{1-\varepsilon_b}{\varepsilon_b} a_p M_i N_i \) | Species mass balance                                                      |
| \( \frac{\partial T}{\partial t} = -\rho C_p u \frac{\partial T}{\partial z} + \frac{\partial}{\partial z} \left( \frac{\partial T}{\partial z} \right) + \frac{4U(T_w - T)}{d_i} \) | Overall energy balance                                                    |
| PM = \( \rho RT \)                           | Equation of state                                                          |
Table 2  Reaction rate equations

| Reaction                                      | Rate Equation                                                                 |
|----------------------------------------------|-------------------------------------------------------------------------------|
| Methanol synthesis from CO (Graaf et al.)    | $r_{\text{CH}_3\text{OH,1}} = \frac{k_1K_{\text{CO}}\left(\varphi_{\text{CO}}\varphi_{\text{H}_2}/\varphi_{\text{CH}_3\text{OH}}\right)^{3/2}}{(1 + K_{\text{CO}}\varphi_{\text{CO}} + K_{\text{CO}}\varphi_{\text{CO}})^{1/2} + \left(\theta^2_{\text{H}_2}/K_{\text{H}_2}\right)}^{1/2}K_{\text{pt}}}$ |
| Water–gas shift (Graaf et al.)               | $r_{\text{CO}} = \frac{k_2K_{\text{CO}}\left(\varphi_{\text{CO}}\varphi_{\text{H}_2}/\varphi_{\text{CH}_3\text{OH}}\right)^{3/2}}{(1 + K_{\text{CO}}\varphi_{\text{CO}} + K_{\text{CO}}\varphi_{\text{CO}})^{1/2} + \left(\theta^2_{\text{H}_2}/K_{\text{H}_2}\right)}^{1/2}K_{\text{pt}}}$ |
| Methanol synthesis from CO$_2$ (Graaf et al.) | $r_{\text{CH}_3\text{OH,2}} = \frac{k_1K_{\text{CO}}\left(\varphi_{\text{CO}}\varphi_{\text{H}_2}/\varphi_{\text{CH}_3\text{OH}}\right)^{3/2}}{(1 + K_{\text{CO}}\varphi_{\text{CO}} + K_{\text{CO}}\varphi_{\text{CO}})^{1/2} + \left(\theta^2_{\text{H}_2}/K_{\text{H}_2}\right)}^{1/2}K_{\text{pt}}}$ |
| Methanol dehydration (Berčič et al.)         | $r_{\text{DME}} = \frac{k_2K_{\text{CH}_3\text{OH}}^{2}\left(c_{\text{CH}_3\text{OH}} - c_{\text{H}_2\text{O}}\text{DME}/K_{\text{pt}}\right)^{3/2}}{1 + 2(K_{\text{CH}_3\text{OH}}c_{\text{CH}_3\text{OH}})^{1/2} + K_{\text{H}_2\text{O}}c_{\text{H}_2\text{O}}}$ |

Fig. 3  Working capacity versus steam partial pressure based on breakthrough experiments for zeolite 3A at 200 °C (red diamonds) and 250 °C (green diamonds), the isotherm prediction in this work (solid lines) and the isotherm prediction fromSips (dashed lines).

Steam adsorbent

In contrast to typical conditions for water adsorption, sorption enhanced DME synthesis requires good sorbent material performance at elevated temperature and pressure. Based on available open literature, selected adsorbents were tested. Next to the physically adsorbing zeolite 3A, 4A and 13X samples, three chemical adsorbents were also tested: LHMC, HHMC, and hydrotalcite. Their performance in terms of the cyclic working capacity for steam adsorption was disregarded for sorption enhanced DME synthesis.

For the most promising SEDMES adsorbent, LTA zeolite 3A, a working isotherm is determined (Fig. 3). In accordance with the open literature about water adsorption on zeolite 3A at lower temperature and pressure, the data can be described with a Sips isotherm. The isotherm parameters are reported in Table 3. As shown in Fig. 3, the Sips isotherm gives an improved prediction of the working capacity, especially at higher steam partial pressure, over the previously used Langmuir–Freundlich isotherm with half the number of parameters.

$$q = q_s \left(\frac{bP}{1 + (bP)^{1/n}}\right)$$  \hspace{1cm} (17)

$$b = b_0 e^{-K_s/RT}$$  \hspace{1cm} (18)

Catalyst performance

The copper–zinc oxide–alumina (CZA) methanol synthesis catalyst is tested separately, and in combination with the methanol dehydration catalyst γ-alumina. As shown in Fig. 4, clearly the DME production is far from equilibrium under direct DME synthesis conditions, whereas the methanol production is close to equilibrium for methanol synthesis (greater than equilibrium in DME synthesis) at temperatures over 250 °C. The optimum temperature for methanol synthesis shifts by dilution of the CZA catalyst from 230 to 250 °C, which is well aligned with temperatures reported for methanol and direct DME synthesis in literature. DME synthesis, however, is far from equilibrium for all catalyst compositions and the DME yield keeps increasing with temperature. Despite the fact that the temperature for methanol dehydration is generally higher, direct DME synthesis is often performed at temperatures of around 250 °C, not only because the methanol synthesis is considered to be the rate determining step in direct DME.

Table 3  Parameters for the Sips isotherm

| Parameter                  | Value |
|----------------------------|-------|
| $q_s$ (kg kg$^{-1}$)       | 0.21  |
| $b_0$ (bar$^{-1}$)         | 4.10 10$^{-7}$ |
| $K_s$ (kJ mol$^{-1}$)      | -60.1 |
| $n$ (-)                    | 1.53  |
larger alcohol–water clusters, but also the (reversible) formation of (surface) boehmite was shown.\textsuperscript{27} Despite the large attention for more active low-temperature methanol dehydration catalysts,\textsuperscript{4,6,59,69,71,73,74,76–79} $\gamma$-Al$_2$O$_3$ remains the catalyst of choice for industrial DME production, due to its low cost, high surface area, good thermal and mechanical stability, and high selectivity to DME because its relatively weak Lewis acid sites do not promote side reactions.\textsuperscript{4,70,76,77}

In contrast to direct DME synthesis, SEDMES offers two specific advantages for the ($\gamma$-Al$_2$O$_3$) catalyst: the system is operated at low steam pressures and is periodically regenerated due to its adsorptive nature. In fact, the reduced steam content will likely promote deactivation (coking) of other, more acidic, dehydration catalysts.\textsuperscript{59,76}

Reaction kinetics for the used materials are determined by fitting the parameters in the methanol synthesis and dehydration reaction models from Graaf \textit{et al.} (1988) and Berčič \textit{et al.} (1992) respectively.\textsuperscript{26,30,31} The new parameters are given in Table 4 and their good predictive capability of the observed concentrations is shown in Fig. 5. For methanol synthesis the average deviation is less than 5\%, which is similar for the DME concentration. Only the methanol concentration in the dehydration experiments shows a higher deviation (11\%) due to increased experimental error by feeding liquid methanol at varying conditions.\textsuperscript{27} The activation energies determined for the methanol synthesis kinetics are lower than the values originally reported, especially considering the conversion of CO, resulting from a higher activity catalyst.\textsuperscript{30} This corresponds with activity factors larger than one reported for present-day methanol synthesis catalysts.\textsuperscript{28,67} Also a difference in catalyst activity for methanol dehydration is observed, where the activation energy is changed 24\% compared to the original value.\textsuperscript{31} The altered value aligns well with modifications of the methanol dehydration kinetics reported for direct DME synthesis in the open literature.\textsuperscript{5}

### Sorption enhanced DME synthesis

Proof-of-concept for sorption enhanced DME synthesis is demonstrated experimentally at a 2 litre bench-scale reactor (TRL4), already a large step forward in the development of the SEDMES process.\textsuperscript{8} Fig. 6 shows a representative

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**Table 4** Model parameters for methanol synthesis and dehydration reaction kinetics\textsuperscript{26}

| Parameter | Value (kJ mol$^{-1}$) | Deviation from original |
|-----------|-----------------------|-------------------------|
| $E_a$ ($k_1$) | 68.1 | -38\% |
| $E_a$ ($k_2$) | 107 | -13\% |
| $E_a$ ($k_3$) | 54.3 | -17\% |
| $AH$ ($k_{C2}$) | -15.7 | -73\% |
| $AH$ ($k_{C3}$) | -56.0 | -17\% |
| $AH$ ($k_{H2,0}/k_{H2}^{1/2}$) | -107 | +2.4\% |
| $E_a$ ($k_4$) | 109 | -24\% |
| $AH$ ($k_{C3,H,OH}$) | -69.6 | -1.3\% |
| $AH$ ($k_{H2,0}$) | -39.3 | -4.4\% |
breakthrough experiment of sorption enhanced DME synthesis. Prior to steam breakthrough, DME and unconverted CO are the primary products. After steam breakthrough the concentration of DME drops, accompanied by the breakthrough of CO₂ and methanol indicating saturation of the adsorbent. As can be seen in Fig. 6 as well, the dynamic cycle model, using the reaction kinetics and water adsorption isotherm as determined in the previous sections, describes the experimental concentration and dynamic behaviour well.

Based on previous modelling work, it was concluded that pressure swing regeneration rather than the so-far required time and energy intensive temperature swing regeneration would be effective. Evaluation of regeneration strategies including pressure swing was therefore among the aims of testing under industrially relevant conditions. The testing under industrially relevant conditions performed in this work indeed proves high performance with pressure swing regeneration, demonstrating over 80% integral carbon selectivity towards DME when using pressure swing regeneration, without the need for a temperature swing (Fig. 6). Where a similar conversion and selectivity can be obtained with pressure swing regeneration (PSA) in comparison with previously reported experiments with a combined temperature and pressure swing regeneration (TPSA), the faster pressure swing regeneration already increases the DME productivity by a factor four, with even further optimisation possible.

One of the optimisation parameters is the carbon selectivity towards DME. Thermodynamically the carbon selectivity to DME is unfavourable and CO₂ will be the main carbon containing product (Fig. 7). However, sorption enhanced DME synthesis allows for a high single-pass carbon conversion to DME irrespective of the carbon source (CO or CO₂), 80% shown here. The model prediction for a CO₂:CO feed of 2:1 is very good (Fig. 7a), and despite a small overprediction for a CO₂ feed (Fig. 7b), the model prediction is still adequate. This overprediction is caused by an apparent catalyst deactivation during the initial part of the experimental campaign, which stabilizes over the full length of the campaign, as shown in Fig. 8. It is well known that especially the CZA catalyst is prone to deactivation under more severe hydrothermal conditions. An advantage of the sorption enhanced reaction conditions include the extremely low water concentration, protecting the catalyst from hydrothermal sintering. However, water has also shown to have a positive influence in catalyst deactivation (by coking) for a CO-rich feed. The cause of the observed small decrease in catalyst activity is subject of follow-up work.

Carbon selectivity and productivity

Rather than just a given value, the carbon selectivity can be chosen and optimised by determining the relative time of the reactive adsorption step due to the dynamic nature of sorption enhanced processes. Fig. 9 shows the change in
carbon selectivity for the carbon containing species with respect to the adsorption time. Besides optimisation by operating conditions as for conventional processes, the dynamic nature of SEDMES allows extra degrees of freedom and therefore, additional flexibility. Experimental carbon selectivity is reported (in other figures) as the integrated selectivity until 5% CO₂ is observed, loosely based on the desired high single-pass conversion of SEDMES and the reduced downstream purification requirements.

The gas hourly space velocity (GHSV) of the feed imposes a trade-off between productivity and selectivity, which is shown in Fig. 10. Regarding this trade-off, the optimal GHSV would be as low as possible from a selectivity point-of-view. Although mass and heat transport eventually would affect the selectivity as well. With increasing GHSV the productivity increases with a loss in selectivity, until the selectivity loss becomes dominant and the productivity will drop as well. In contrast to conventional “steady-state” reaction conditions, the SEDMES process has extra degrees of freedom to optimise the GHSV in combination with the cycle design and timing of the sorption enhanced reaction steps. The selectivity and productivity need to be balanced in the process design and techno-economic evaluation for a specific case.

The duration of the regenerative purge step is one of these additional parameters. A longer purge time results in better regeneration of the system and therefore a higher DME selectivity (Fig. 11), as seen experimentally and well predicted by the model. A longer purge time relative to the adsorption time, however, would require a cycle design with more columns, resulting in a lower overall specific productivity (kg h⁻¹ m⁻³). To decrease the inventory (m³) and therefore increase the overall productivity a short purge time would be desired. Fig. 11 shows a discrepancy between experimental results and model predictions for a 30 minute purge time. This can be explained by the notion that the experimental data points show a decreasing trend (rather than a spread as is the case for the other data) towards the modelled selectivity.

Fig. 12 shows the experimental results and model predictions accordingly for various combinations of the
adsorption and regeneration time. It must be noted that the experimental results are based on 2–4 point integration for short timings. Nonetheless, the model predicts the experimental results well for lowering the adsorption and purge time to 20/30 and 10/20 minutes respectively. While the model predicts a significant drop in selectivity for 10/10 minutes due to a decreasing working capacity of the adsorbent, the experimental results show a significantly smaller drop. Investigation of the adsorption behaviour during faster cycling is required to further clarify these observations.

The combination of the adsorption duration (ADS) and regeneration time (DES) allows optimising the trade-off for DME selectivity and productivity, as shown in Fig. 13. Shorter adsorption times potentially result in an increased production rate. The larger reactor column requirement when the purge time does not decrease with the adsorption time, however, results in a drop in cyclic productivity (kg h⁻¹ m⁻³). Looking at the minimum number of columns required for any given adsorption and purge time, the productivity could be significantly boosted for shorter cycle times with the highest ADS/DES ratio (red bars in Fig. 13). The loss in carbon selectivity predicted by the model for 10/10 minutes would also result in a major drop in predicted productivity (Fig. 12). The promising experimental results, shown in Fig. 13, however indicate that a minor loss in selectivity could still result in increased productivity for faster cycling. The productivities reported in Fig. 13 correspond to 0.04–0.06 kg h⁻¹ kgcat⁻¹. This is a major improvement to the previously reported TPSA cycle and close to direct DME pilot plant.
For the first time, a validated pressure swing regeneration cycle for sorption enhanced DME synthesis (SEDMES) is demonstrated under industrially relevant conditions. SEDMES is a highly flexible process for converting CO$_2$-rich (bio-based) syngas and CO$_2$ directly to DME with a high single-pass conversion, reducing or even eliminating the conventional large recycles and downstream purification sections. The industrially relevant testing performed in this work indeed proves this significant performance, 80% single-pass carbon selectivity to DME demonstrated with pressure swing regeneration, which already allows for a factor four increase in productivity with further optimisation still possible.

The study of the adsorbent material, the catalysts, and the combination of all materials involved resulted in a validated dynamic reactor model, which allows adequate upscaling of the SEDMES technology and predictions of large scale DME synthesis for which faster cycling seems promising to further enhance productivity. Techno-economic and life cycle analyses have to be performed to investigate the economic and carbon mitigating benefits of the high efficiency carbon utilisation technology.

4. Conclusions

For the first time, a validated pressure swing regeneration cycle for sorption enhanced DME synthesis (SEDMES) is demonstrated under industrially relevant conditions. SEDMES is a highly flexible process for converting CO$_2$-rich feed, which would strongly deteriorate productivity for CO to DME, to DME, CO, CO$_2$, and methanol with a high single-pass conversion, reducing or even eliminating the conventional large recycles and downstream purification sections. The industrially relevant testing performed in this work indeed proves this significant performance, 80% single-pass carbon selectivity to DME demonstrated with pressure swing regeneration, which already allows for a factor four increase in productivity with further optimisation still possible.

The study of the adsorbent material, the catalysts, and the combination of all materials involved resulted in a validated dynamic reactor model, which allows adequate upscaling of the SEDMES technology and predictions of large scale DME synthesis for which faster cycling seems promising to further enhance productivity. Techno-economic and life cycle analyses have to be performed to investigate the economic and carbon mitigating benefits of the high efficiency carbon utilisation technology.

**Nomenclature**

- $a_p$: Particle interfacial area ($m^2 m^{-3}$)
- $b$: Isotherm equilibrium constant (bar$^{-1}$)
- $c_i$: Concentration of component $i$ (mol m$^{-3}$)
- $C_p$: Gas thermal conductivity ($J kg^{-1} K^{-1}$)
- $C_{p,p}$: Particle thermal conductivity ($J kg^{-1} K^{-1}$)
- $d_p$: Particle diameter ($m$)
- $D_a$: Axial dispersion coefficient ($m^2 s^{-1}$)
- $E_a$: Activation energy ($kJ mol^{-1}$)
- $G$: Ergun constant ($-$)
- $\Delta H_{ads}$: Adsorption enthalpy ($J mol^{-1}$)
- $\Delta H_{f,i}$: Reaction enthalpy ($J mol^{-1}$)
- $k$: Reaction rate constant (mol s$^{-1}$ kg$^{-1}$ bar$^{-1}$) or (kmol kg$^{-1}$ h$^{-1}$)
- $K_i$: Adsorption equilibrium constant of component $i$ (bar$^{-1}$) or (m$^3$ kmol$^{-1}$)
- $K_p$: Equilibrium constant (based on partial pressure) ($-$)
- $M_i$: Molecular weight of component $i$ (kg mol$^{-1}$)
- $N_i$: Mole flux of component $i$ (mol m$^{-2}$ s$^{-1}$)
- $P$: Reactor pressure (bara)
- $P_i$: Partial pressure of component $i$ (bara)
- $q_i$: Adsorbent loading (mol kg$^{-1}$)
- $q_s$: Saturation capacity (kg$^{-1}$)
- $r_i$: Reaction rate of component $i$ (mol m$^{-3}$ s$^{-1}$) or (mol kg$^{-1}$ s$^{-1}$) or (kmol kg$^{-1}$ h$^{-1}$)
- $R$: Ideal gas constant ($J mol^{-1} K^{-1}$)
- $t$: Time (s)
- $T$: Temperature (K)
- $u$: Superficial gas velocity ($m s^{-1}$)
- $U$: Overall heat transfer coefficient ($W m^{-2} K^{-1}$)
- $v$: Interstitial gas velocity ($m s^{-1}$)
- $z$: Axial coordinate ($m$)

**Greek letters**

- $\varepsilon_b$: Bed voidage ($-$)
\( \lambda \) Axial thermal conductivity (W m\(^{-1}\) K\(^{-1}\))

\( \rho \) Density (kg m\(^{-3}\))

\( \rho_p \) Particle density (kg m\(^{-3}\))

\( \phi_i \) Partial fugacity of component i (bara)

\( \omega_i \) Weight fraction of component i (~)

Conflicts of interest

There are no conflicts to declare.

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