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A Novel Approach for Kinetic Measurements in Exothermic Fixed Bed Reactors: Advancements in Non-Isothermal Bed Conditions Demonstrated for Methanol Synthesis

F. Nestler, V. P. Müller, M. Ouda, M. J. Hadrich, A. Schaad, S. Bajohr and T. Kolb

Kinetic modelling of methanol synthesis remains one key challenge for the implementation of Power-to-Methanol technologies based on CO₂-rich gas streams and sustainably produced H₂. Within this work, a novel approach for kinetic model validation and parameter estimation using an experimental miniplant setup with polytropic bed conditions is presented. The miniplant setup features a highly resolved fibre optic temperature measurement combined with FTIR product composition analysis. Comparison of the experimental temperature and concentration data to a simulation model applying literature kinetic models, confirmed the necessity of axial experimental data to deliver an appropriate kinetic description of the methanol synthesis reaction network. A refitting of the literature kinetic models was performed in order to enhance their capability to account for the catalytic behaviour of a modern commercial catalyst. Besides the traditional measurement of the outlet concentration, it was shown that the temperature profile as a direct consequence of exothermic reactions in polytropic miniplant setups can be used to derive an improved kinetic description if appropriate models for heat transfer and diffusion are provided. Finally, the behaviour of the proposed new kinetic model is discussed on the industrial scale by means of a sensitivity analysis emphasizing the applicability of the presented novel approach for the scale-up from miniplant to industrial scale.

Introduction

The rising demand for energy carriers and base chemicals produced from sustainably generated hydrogen (H₂), e.g. by water electrolysis, in the context of Power-to-X (PtX) processes recently created vast research activity with the aim of making alternative synthesis routes competitive to their fossil counterpart. Among the most promising PtX products, ammonia (NH₃), synthetic natural gas (SNG) and methanol (MeOH) are discussed. In this context, methanol is very likely to play a key role due to its capability as carbon dioxide (CO₂)-sink when combined with industrial processes, as cement or steel production. Moreover, methanol has the advantage of an already existing trade infrastructure and is a key molecule for production of high value derivatives such as polymers, fuels and olefins. With an annual production capacity of approximately 100 Mt methanol is already today an important platform molecule for the chemical industry and the energy sector.

Despite the fact of methanol synthesis being one of the oldest thermochemical high pressure processes, questions remain open on the adaption of the process from fossil-based synthesis gas (syngas) with high carbon monoxide (CO)-contents towards sustainable syngas with high CO₂-contents. As both, electrolytically produced H₂ from renewable energy and carbon dioxide-rich gas obtained from the coupled industrial process are subjected to fluctuations, dynamic description of the methanol synthesis process and the synthesis reactor are imperative for the implementation of PtM processes. However, dynamic operation of the methanol synthesis reactor demands for a validated simulation including a highly reliable kinetic model. Therefore, an improved kinetic understanding of methanol synthesis is one key issue for the implementation of PtM technology on the industrial scale.

In general, methanol synthesis carried out on commercial Cu/Zn/Al₂O₃-catalysts can be expressed via the following exothermic equilibrium limited reactions:

\[
\begin{align*}
\text{CO}_2(g) + 3\text{H}_2(g) &\rightleftharpoons \text{CH}_3\text{OH}(g) + \text{H}_2\text{O}(g) \quad \Delta H^\circ = -50 \text{kJ mol}^{-1} \quad (1) \\
\text{CO}(g) + \text{H}_2\text{O}(g) &\rightleftharpoons \text{CO}_2(g) + \text{H}_2(g) \quad \Delta H^\circ = -41 \text{kJ mol}^{-1} \quad (2) \\
2\text{CO}(g) + 2\text{H}_2(g) &\rightleftharpoons \text{CH}_3\text{O}_2H(g) \quad \Delta H^\circ = -91 \text{kJ mol}^{-1} \quad (3)
\end{align*}
\]

In this reaction network, direct CO-hydrogenation (Eq. (3)) was proven to proceed only to a limited extent but rather as a combination of CO₂-hydrogenation (Eq. (1)) and water-gas-shift-reaction (WGS, Eq. (2)). Various modelling approaches describing methanol synthesis kinetics can be found in...
literature\textsuperscript{13–20}, however, the mechanistic description remains a controversial topic in the scientific community\textsuperscript{20}. For reactor design, the temperature and the axial position of the hot spot, i.e. the point with the highest temperature inside the reactor, represent important key parameters\textsuperscript{21–26}. In one of our previous studies we demonstrated significant discrepancies regarding the simulative description of the hot spot depending on the choice of the kinetic model\textsuperscript{27}. Consequently, substantial uncertainties in reactor and process design are obtained leading towards oversizing of process equipment and application of too mild process conditions, e.g. low synthesis temperatures\textsuperscript{22}. Classic kinetic experimental setups are built with respect to minimal temperature gradients along the radial and axial coordinate of the fixed bed allowing a simple temperature measurement and straightforward validation of reaction kinetics\textsuperscript{28–30}. Besides pressure and temperature, key parameters in kinetic measurements in methanol synthesis usually address the variation of the carbon oxide ratio (COR), the stoichiometric number (SN) and the gas hourly space velocity (GHSV). The three parameters are formulated as follows\textsuperscript{4}:

\[
\text{GHSV} = \frac{V_{\text{norm}}}{V_{\text{cat}}} \tag{4}
\]

\[
\text{COR} = \frac{y_{\text{CO}}}{y_{\text{CO}_2} + y_{\text{CO}}} \tag{5}
\]

\[
\text{SN} = \frac{y_{\text{H}_2} - y_{\text{CO}_2}}{y_{\text{CO}_2} + y_{\text{CO}}} \tag{6}
\]

In order to reduce the risks of a scale-up from lab scale to the industrial scale, a wide range of operating conditions needs to be covered in classic kinetic measurements. However, as the composition of the products at the reactor outlet is usually the only information gathered by these experiments, the kinetic models derived often rely on limited validation data\textsuperscript{25–17,20}. Due to the lack of experimental data, possible issues arise when the kinetic models derived in ideally isothermal fixed bed reactors are transferred to an industrial reactor featuring non-isothermal bed conditions. Differentially resolved kinetic data would be mandatory in order to deliver a local reaction rate along the axial reactor dimension in an industrial reactor with appropriate accuracy\textsuperscript{27}

This work introduces a novel approach applying a polytropic miniplant reactor scaled down from an industrial steam cooled multi-tubular reactor for the validation of kinetic models for exothermic fixed bed reactions. The validation relies on a highly resolved measurement of the axial temperature profile inside the reactor in addition to the product gas analysis. By gathering information on the axial temperature profile, the quantity of validation information can be increased significantly in comparison to traditional integral fixed bed measurements. Importantly, the transferability of the derived improved kinetic model on industrial scale will be demonstrated. Moreover, an optimization of the miniplant dimensions will be carried out in order to increase the similarities between industrial and miniplant reactor scale.

\textbf{Methods}

The validation approach presented within this work strongly relies on a detailed simulation model of the miniplant reactor. In order to discuss the derived kinetic model also on the industrial scale, ability of the simulation to adapt to this scale was of high importance for this work. Therefore, the implementation of the simulation platform used in this work will be described first. Subsequently, the scale-down and design of the experimental setup as well as the methodology for the kinetic parameter estimation will be presented. Finally, the methodology for the discussion of the derived kinetic model on the industrial scale will be explained.

\textbf{Simulation platform}

In order to use a non-isothermal experimental setup for validation and parameter fitting of a kinetic model, a simulation platform for the description of the methanol synthesis reactor was developed. This platform can be described as a wrapper allowing the reactor simulation to be adaptable to different scales and geometries. The reactor model is based on sub-models describing heat transfer, powder kinetics and diffusion inside the reactor. As shown in Figure 1, powder kinetics and diffusion model express the effective particle kinetics, which are then included into the numerical one-dimensional reactor model.

\textbf{Heat transfer.} The sub-model accounting for the heat transfer between reaction zone and cooling medium is based on VDI Heat Atlas providing established state-of-the-art approaches for engineering heat transfer problems\textsuperscript{31}. This sub-model provides semi-empirical approaches for heat transfer in packed beds with gas flow ($\lambda_{\text{rad}}$, $\alpha_{\text{int}}$) and the heat transfer outside the reactor tube ($\alpha_{\text{ext}}$). In case of the miniplant $\alpha_{\text{int}}$ was expressed by convective thermal oil heat transfer, while for the industrial scale a steam cooling was applied, respectively\textsuperscript{31}. Overall heat transfer coefficient $U$ was calculated as follows\textsuperscript{32}:

\[
U = \frac{1}{\frac{1}{\alpha_{\text{int}}} + \frac{d_{\text{int}}}{8 \cdot \lambda_{\text{rad}}} + \frac{d_{\text{int}} \cdot \ln \left( \frac{d_{\text{ext}}}{d_{\text{int}}} \right)}{2 \cdot \lambda_{\text{wall}}} + \frac{d_{\text{int}}}{\alpha_{\text{ext}} \cdot d_{\text{ext}}}} \tag{7}
\]

Figure 1: Simulation platform applied within this publication.
The effective thermal conductivity of the catalyst particles was set to 0.33 W m⁻¹ K⁻¹ according to Henkel[33], who performed a comprehensive analysis of the thermophysical and diffusional properties of a comparable catalyst in his PhD work.  

**Powder Kinetics.** Some of the most commonly used kinetic models for methanol synthesis are the models provided by Graaf[34–37] and Bussche-Froment[38–41]. Graaf’s kinetic model is based on the stepwise hydration of CO and CO₂ with the rate determining step (rds) determined by an error discussion of the 48 possible combinations of rds of CO-hydrogenation, CO₂-hydrogenation and reverse WGS (rWGS) [13,42,43]. In contrast to the mechanism applied by Graaf, Bussche and Froment considered a different reaction mechanism based only on CO₂-hydrogenation coupled with rWGS via the formyl species[35]. However, in our previous work inconsistencies of both kinetic models were proven by means of a comprehensive computational simulation study[37]. In this previous work we also proposed a new kinetic model based on measured data published in scientific literature[33] and Graaf’s kinetic approach, though with CO₂-hydrogenation removed[33]. The elementary steps of the reaction network can be expressed as follows:

**CO₂-hydrogenation**

\[
\begin{align*}
A1 & : \text{CO}_2{s1} + H^* {s2} \rightarrow \text{HCO}_2{s1} + s2 & (8) \\
A2 & : \text{HCO}_2{s1} + H^* {s2} \rightarrow \text{H}_2\text{CO}_2{s1} + s2 & (9) \\
A3 & : \text{H}_2\text{CO}_2{s1} + H^* {s2} \rightarrow \text{HCO}_2{s1} + \text{H}_2\text{O}{s2} + s2 & (10) \\
A4 & : \text{H}_2\text{CO}_2{s1} + \text{H}_2\text{O}{s2} \rightarrow \text{HCO}_2{s1} + \text{H}_2\text{O}{s2} + s2 & (11) \\
A5 & : \text{H}_2\text{CO}_2{s1} + \text{H}_2\text{O}{s2} \rightarrow \text{H}_2\text{CO}_2{s1} + \text{H}_2\text{O}{s2} + s2 & (12) \\
A6 & : \text{H}_2\text{CO}_2{s1} + \text{H}_2\text{O}{s2} \rightarrow \text{CH}_3\text{OH} + s1 + s2 & (13) \\
\end{align*}
\]

**rWGS**

\[
\begin{align*}
B1 & : \text{CO}_2{s1} + H^* {s2} \rightarrow \text{HCO}_2{s1} + s2 & (14) \\
B2 & : \text{HCO}_2{s1} + \text{H}_2\text{O}{s2} \rightarrow \text{CO}_2{s1} + \text{H}_2\text{O}{s2} + s2 & (15) \\
\end{align*}
\]

In agreement to Graaf’s finding, the steps A3 and B2 were considered rate determining. The rate equations for all other possible combinations of CO₂-hydrogenation and rWGS were derived and fitted to the experimental data obtained in this work regarding to the procedure introduced later in this chapter. However, as the combination A3B2 showed the lowest remaining errors between the experimental data obtained in this study and the simulation among all the possible combinations of Eq. (8) to Eq. (15), these were not further considered within this study. Consequently, the rate equations for the kinetic model which was already applied in our previous study can be formulated as follows[37,38]:

\[
\begin{align*}
r_{\text{CO}_2} &= \frac{k_1 \cdot k_2 \cdot f_{\text{CO}_2} \cdot f_{\text{H}_2}\text{O}}{(1 + K_1 \cdot f_{\text{CO}} + K_2 \cdot f_{\text{CO}_2} + K_3 \cdot f_{\text{H}_2}\text{O})} \cdot \text{EQ1} & (16) \\
r_{\text{rWGS}} &= \frac{k_2 \cdot k_2 \cdot f_{\text{CO}_2} \cdot f_{\text{H}_2}\text{O} \cdot \text{EQ2}}{(1 + K_1 \cdot f_{\text{CO}} + K_2 \cdot f_{\text{CO}_2} + K_3 \cdot f_{\text{H}_2}\text{O})} & (17) \\
\end{align*}
\]

The equilibrium terms for CO₂-hydrogenation (EQ₁) and rWGS (EQ₂) can be expressed as follows according to Graaf et al.[44]:

\[
\begin{align*}
\text{EQ1} &= 1 - \frac{f_{\text{CH}_3\text{OH}} \cdot f_{\text{H}_2}\text{O}}{f_{\text{CO}_2} \cdot f_{\text{H}_2}\text{O} \cdot K_{\text{eq1}}} & (18) \\
\text{EQ2} &= 1 - \frac{f_{\text{CO}_2} \cdot f_{\text{H}_2}\text{O} \cdot K_{\text{eq2}}}{f_{\text{CO}_2} \cdot f_{\text{H}_2}\text{O} \cdot K_{\text{eq2}}} & (19) \\
\end{align*}
\]

The fugacities were calculated using the Soave-Redlich-Kwong equation of state (SRK EoS)[65]. The model parameters of the above mentioned kinetic models are expressed by an Arrhenius correlation as follows for the kinetic rate constants \(k_i\) and adsorption constants \(K_i\):

\[
\begin{align*}
k_i &= A_ki \cdot \exp \left(- \frac{B_ki}{R \cdot T} \right) & (20) \\
K_i &= A_ki \cdot \exp \left(- \frac{B_ki}{R \cdot T} \right) & (21) \\
\end{align*}
\]

The semi-empirical constants \(A_ki\) and \(B_ki\), as well as \(A_ki\) and \(B_ki\), in Eq. (20) and (21), respectively, can be tuned in order to obtain agreement between experimental data and reactor simulation[46]. Further details on the kinetic models of Graaf and Bussche are provided in our previous work[27] and in the supplementary material of this work.

**Diffusion.** Methanol synthesis is known to be subjected to mass transfer limitations depending on particle size and reaction conditions applied[47,48]. The Thiele modulus\[49\] \(\Phi_M\) is frequently applied in scientific literature to describe the diffusion limitation caused by the reactants passing through the porous structure of the catalyst towards the active sites\[40,50,51\]. Calculation methodology was adopted from Lommerts et al.[50] who discussed the applicability of Thiele modulus against the Dusty Gas Model: In their study the authors stated Thiele modulus as an appropriate compromise between computational time and accuracy. Thiele modulus was calculated via the pseudo-first order reaction velocity with \(k_{\text{eq,CO}_2}\) and \(k_{\text{eq,Methanol}}\) representing the pseudo equilibrium constant[50] as well as \(k_{\text{u,CO}_2}\) and \(k_{\text{u,Methanol}}\) as pseudo-first-order rate constant[42]:

\[
\Phi_M = \frac{d_p}{6} \frac{k_{\text{u,CO}_2} \cdot (K'_{\text{eq,CO}_2} + 1)}{D_{\text{m,CO}_2} \cdot k_{\text{u,Methanol}} \cdot k_{\text{eq,Methanol}}} & (22)
\]

Effective diffusion coefficients for water and methanol in the mixture were obtained as follows[50,52]:

\[
\frac{1}{D_{\text{m,CO}_2}} = \frac{\tau \cdot \sum_{j=1}^{N} \frac{1}{D_{i,j}} + \frac{1}{D_{K,j}}}{\sum_{j=1}^{N} \frac{1}{D_{i,j}} + \frac{1}{D_{K,j}}} & (23)
\]

Diffusion coefficient for the single component in the reaction mixture \(D_{ij}\) was calculated according to Fuller et al.[53] while calculation of Knudsen diffusion was performed with regard to Westerterp et al.[52]. Tortuosity \(\tau\) and porosity of the catalyst \(\varepsilon_p\) were chosen with respect to Henkel[33] (values see Table 1).

| Parameter | Unit | Value |
|-----------|------|-------|
| \(\tau\)  |      | 2.99  |
| \(\varepsilon_p\)|   | 0.58  |

The efficiency factor of the reactions \(\eta_{eff}\) is calculated for both, water and methanol as follows[50,52]:

\[
\eta_{eff} = \frac{1}{1 - \frac{1}{D_{\text{m,CO}_2} \cdot k_{\text{u,Methanol}} \cdot k_{\text{eq,Methanol}}}} & (24)
\]

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Amongst the two efficiency factors obtained for water and methanol, the smaller value was considered for the effective reaction rate in order to describe the maximum overall diffusion limitation in the reaction network.

**Steady state reactor model.** Based on the sub-models for heat transfer, powder kinetics and diffusion, a steady state reactor model can be built utilising the following differential equations for mass, energy, and momentum balance:

\[
\frac{dh_j}{dx} = \rho_{\text{bulk}} \cdot A_R \cdot \sum u_j \cdot \tau_{\text{eff},j}
\]

\[
\frac{dT}{dx} = \frac{\Delta H_{R,j}}{c_p, f} \cdot \tau_{\text{eff},j} \cdot A_R \cdot \rho_{\text{bulk}} \cdot \sum u_j \cdot \tau_{\text{eff},j} + \frac{\pi \cdot \rho_{\text{int}} \cdot U \cdot (T_{\text{cool}} - T)}{c_p, f \cdot r_{\text{tot}}}
\]

\[
\frac{dp}{dx} = - \left( 1.75 + 150 \cdot \frac{1 - \varepsilon_{\text{bulk}}}{R e_p} \right) \cdot \frac{1 - \varepsilon_{\text{bulk}}}{\varepsilon_{\text{bulk}}^3 \cdot d_p} \cdot \rho_f \cdot u_0^2
\]

Reynolds particle number \(Re_p\) applied in Erguns Equation (Eq. (27)) was calculated as follows:

\[Re_p = \frac{u_0 \cdot d_p}{\nu_f}\]

Empty tube fluid velocity \(u_0\) in the reactor was calculated by continuity equation applying the empty tube cross section \(A_R\) and the volumetric flow rate \(V\) at rated pressure and temperature:

\[u_0 = \frac{V}{A_R}\]

Thermophysical properties of the gas phase as heat capacity \(c_p, p\) and kinematic viscosity \(\nu\) were calculated according to DIPPR equations\(^{54}\), SRK EoS\(^{55}\) and VDI Heat Atlas\(^{31}\). Due to the nature of the one-dimensional model, radial gradients inside the reactor were neglected in the simulation. However, the effect of this assumption towards the parameter fitting and scale-up could be investigated in future studies, utilizing more powerful computational resources.

**Industrial reactor simulation parameters.** The reactor of Shiraz methanol synthesis plant in Iran was used as a reference for the industrial scale in this study, as it is well documented in scientific literature\(^{37,56-57}\). The steam cooled reactor is part of a conventional methanol synthesis facility producing methanol from a syngas obtained by natural gas reforming\(^{58}\). Design data of the reactor are listed in Table 2.

| Parameter | Unit | Value |
|-----------|------|-------|
| \(d_{\text{int}}\) | m | 3.8 \cdot 10^{-2} |
| \(h_{\text{cat}}\) | m | 7.022 |
| \(d_p\) | m | 5.4 \cdot 10^{-3} |
| \(\varepsilon_{\text{bulk}}\) | - | 0.39 |
| \(\rho_{\text{bulk}}\) | kg m\(^{-3}\) | 1132 |

**Experimental**

**Scale-down of the industrial reactor.** Design of reactors for kinetic measurements is a complex topic subjected by numerous scientific studies\(^{28,58,59}\). In general, dimensions of kinetic setups are determined by dimensionless index number as e.g.:

- The Bodenstein number with the axial dispersion coefficient calculated according to Kraume\(^{60}\)

\[Bo = \frac{u_0 \cdot h_{\text{cat}}}{D_{ax}}\]  (30)

- The Reynolds particle number (see Eq. (28))
- The reactor-particle diameter ratio\(^{61}\).

\[\varphi = \frac{d_{\text{int}}}{d_p}\]  (31)

However, already in 1938 Damköhler et al. found that the scale-down of heterogeneous fixed bed reactors is not possible without violation of the terms of similarity\(^{62}\). Therefore, in classical kinetic setups ideal conditions by means of fluid dynamics, thermal operation and diffusion are acquired, which however, significantly differ from the industrial scale\(^{28}\). Hence, multiple experimental campaigns need to be executed to accurately transfer the results of small-scale measurements towards industrial scale.

With regard to methanol synthesis as one of the oldest high pressure reactions, versatile research has been performed and rich knowledge on the modelling of heat transfer, kinetics and diffusion was published in the past decades\(^{10,63}\). Therefore, a simulation-based approach was realized in our work in order to design an experimental miniplant setup with a high transferability of the experimental results towards industrial scale. As a key parameter, the GHSV was held equal for both industrial and miniplant reactor simulation in order to obtain similar residence times inside the reactor on both scales.

Based on the approach and infrastructural boundary conditions, the miniplant setup used within this work was designed and built. The following procedure was applied to design the reactor dimensions:

1.) Definition of the miniplant scale considering the lab infrastructure

2.) Design of a cooling system for a comparable heat transfer in the miniplant related to the industrial scale

3.) Optimization of the reactor dimensions by minimizing the difference between the simulated temperature profiles of the industrial reactor and the miniplant using the simulation platform

In contrast to the industrial reactor implemented as multi tubular steam cooled reactor, the miniplant reactor consists of a double pipe arrangement with thermal oil circulated through the annular gap while the catalyst is placed inside the inner tube. The idea behind the thermal oil cooling of the miniplant was to counter-balance the higher cooling-area-to-catalyst-volume ratio of the miniplant in comparison to the industrial reactor. The overall objective of this advanced cooling concept was to achieve a temperature profile inside the miniplant reactor comparable to that of the industrial reactor.

In order to determine the reactor dimensions for a maximized comparability of the miniplant setup to the industrial scale, the diameter of the miniplant setup was varied with the catalyst...
bed length adjusted to the previously defined catalyst volume \( V_{\text{cat}} \) correspondingly:

\[
h_{\text{cat}} = \frac{4 \cdot V_{\text{cat}}}{\pi \cdot d_{\text{cat}}^2}
\]

(32)

Temperature profile inside the reactor was identified one key indicator for the similarities between industrial and miniplant reactor. Therefore, the RMSE between the temperature profiles was considered as objective function for the optimization of the miniplant geometry:

\[
RMSE_T = \sqrt{\frac{\sum_{i=1}^{N_{\text{inc}}} (T_{x,i,\text{ind}} - T_{x,i,\text{miniplant}})^2}{N_{\text{inc}}}}
\]

RMSE\(_T\) was minimized by the Nelder Mead algorithm implemented as a fminsearch algorithm in MATLAB\(^6\).

By performing simulations for both, industrial and miniplant setup, the dimensions listed in Table 3 were iteratively defined.

Table 3: Parameters applied for the reactor simulation of the industrial reactor and the miniplant setup.

| Parameter | Unit | Value |
|-----------|------|-------|
| \( d_{\text{cat}} \) | m | 1.3 \times 10^{-2} |
| \( d_{\text{ext}} \) | m | 1.6 \times 10^{-2} |
| \( h_{\text{cat}} \) | m | 1.12' |
| \( d_p \) | m | 1.0 \times 10^{-3}'' |
| \( \rho_{\text{bulk}} \) | kg m\(^{-3}\) | 1134'' |
| \( d_{\text{shell}} \) | m | 2.1 \times 10^{-2}'' |
| \( V_{\text{oil}} \) | l min\(^{-1}\) | 17.1'' |

*\( d_{\text{cat}} \) determined by simulation-based scale-down

**\( d_{\text{cat}} \) determined by experiments

The dimensionless index numbers defined previously are provided for both, industrial and miniplant setup in Table 4.

Table 4: Dimensionless index numbers for industrial and miniplant reactor; Bodenstein number \( Bo \) and Reynold particle number \( Re_p \) were calculated at the following working point: \( T_S = 240 \, ^\circ\text{C}\); \( p = 80 \, \text{bar}\); \( \text{COR} = 0.9\); \( \text{SN} = 4.0\); \( \text{GHSV} = 6,000 \, \text{h}^{-1}\).

| Parameter | Unit | Industrial reactor\(^*\) | Miniplant reactor |
|-----------|------|-------------------------|------------------|
| \( Bo \)  | -    | 1.014                   | 900              |
| \( \varphi \) | -    | 7.04                    | 13               |
| \( Re_p \) | -    | 1185                    | 36               |

Comparison of the dimensionless indices shows that both reactor scales satisfy the criterions for ideal plug flow\(^32\), i.e. \( Bo > 80 \), and non-laminar particle flow, i.e. \( Re_p > 10 \). The difference of the \( Re_p \) numbers between industrial and miniplant reactor due to the adjusted reactor and catalyst geometries was considered in the simulation of pressure loss as well as convective heat transfer inside the reactor. As the reactor-particle-ratio satisfies the criterion \( \varphi > 10 \) wall effects in the miniplant were neglected\(^{30,33}\). However, in case of the industrial reactor \( \varphi \) was below this critical threshold. Measured data obtained from this reactor would be helpful to quantify possible deviations from the herein assumed ideal plug flow behaviour for the industrial reactor.

The inner diameter of 13 mm for the miniplant reactor was obtained by the simulation-based scale-down utilizing the kinetic model by Bussche-Froment\(^{15}\) at high CO\(_2\) contents. As the choice of the kinetic model was found to influence the optimal reactor dimensions significantly, optimized miniplant dimensions applying the kinetic model derived within this work will be presented at the end of this study. These were determined for a wide range of synthesis conditions covering two different pressure levels of 50 bar and 80 bar at \( \text{GHSV} = 9,000 \, \text{h}^{-1}\). Feed gas composition was varied in the range \( 2.0 \leq SN \leq 8.0 \) and \( 0.5 \leq \text{COR} \leq 1.0 \).

**Experimental miniplant setup.** A simplified flow sheet of the miniplant setup utilized within this study is given in Figure 2. The reaction educts CO, CO\(_2\), H\(_2\) as well as the inert gas nitrogen (N\(_2\)) can be flexibly and precisely dosed into the system. Due to the high synthesis pressures up to 80 bar applied to the system, a liquid dosing of CO\(_2\) utilizing an HPLC-pump coupled with a Coriolis flow meter was integrated into the system. The liquid CO\(_2\) was mixed with the other educts and evaporated along a heated line towards the reactor. The correct calibration of the gas dosing unit was frequently verified by a bypass gas phase measurement.

The heat released by the reaction inside the inner tube was removed by thermal oil circulating along the annular gap in counter current flow. Volumetric flow rate of the oil was measured by a rotameter calibrated for the thermal oil used inside the cooling system (Fractoglass X-400-A). Temperature of the thermal oil was controlled by a closed-cycle thermostat.

Figure 2: Simplified flow sheet of the experimental setup.
The reactor was filled with a commercial Cu-based catalyst provided by Clariant. The pelletized catalyst particles were ground and sieved to a particle size of \( d_0 = 1 \) mm to avoid wall effects inside the reactor. An inert bed of \( \alpha \)-alumina supplied by Merck KGaA was placed above and below the catalyst bed. The fixed bed was held inside the reactor with a porous stainless steel support disc. Preliminary tests introducing syngas into the heated reactor filled with only the inert material confirmed the inert behaviour of the whole setup.

In order to gather axial information about the reaction kinetics, the reactor was equipped with a system for fibre optical temperature measurement. The measurement principle of this technology is based on axial variation of the refractive index along a glass fibre due to impurities or local defects. Application of Fourier transformation to a back-scattered light signal leads towards continuous information about (thermal) expansion of the fibre and thereby delivers a spatially resolved temperature information. The glass fibre was placed inside a 0.8 mm steel capillary; the optical signal was generated and processed by a Luna ODiSI 6102 unit. An axial resolution of \( \Delta x = 2.6 \) mm was selected for the measurement campaign leading to 431 measurement increments for temperature measurement \( (N_{\text{meas}}) \) inside the catalyst bed with a length of 1.12 m. Calibration of the fibre was carried out by heating the thermal oil cycle to constant temperature levels between 50 °C and 265 °C. The oil inlet and outlet temperatures were measured by two Pt-100 temperature sensors at the oil inlet (TI02) and outlet (TI01); a heat loss resulting in a temperature decrease of approximately 1.5 K between thermal oil inlet and outlet was regarded by a linear temperature decrease along the reactor. The cooling temperature \( T_{\text{cool}} \) was adjusted to the reading of TI01. As a result of the calibration a polynomial of 3rd degree was determined for each increment along the fibre. Extrapolation of these polynomials was performed for temperatures exceeding 265 °C as the thermal oil did not allow for higher calibration temperatures due reasons of plant safety. For analysis of the reaction products a MKS MultiGas™ 2030 online FTIR with an optical path length of 35 cm was used for quantitative product analysis. Since \( H_2 \) as a homonuclear gas cannot be detected by FTIR, the molar fraction of this gas was determined by the component balance as follows:

\[
y_{\text{H}_2} = 1 - \sum_{i=1}^{N_{\text{comp}}} y_i
\]  
(34)

Besides the main reactants CO, \( \text{CO}_2 \), \( \text{H}_2 \text{O} \) and \( \text{MeOH} \), side products such as methyl formate, methane, ethanol, acetone and acetic acid were calibrated and analysed by the FTIR. However, their low concentrations of less than 100 ppm in the product gas led towards a high signal-to-noise ratio, making exact quantification in the gas phase impossible. Therefore, the side products mentioned above were excluded from the mass balance over the reactor.

Besides the gas phase analysis, the main product stream was led through a cooler-condenser unit at an operating temperature of 10 °C to separate the liquid products from the gas phase for qualitative analysis of condensable trace compounds. Analysis of the liquid phase was carried out using NMR spectroscopy.

All real time information on sensor properties such as volumetric flow rates, inlet and outlet pressures, temperature profile as well as gas phase composition were logged with a sample rate of 1 Hz.

**Experimental plan.** In order to determine the reaction kinetics over a wide parameter range relevant for application with CO2-rich syngas, a comprehensive experimental plan was executed. Besides pressure, the parameters GHSV, COR and SN were varied in the experimental campaign due to their relevance for reaction kinetics. The variation ranges of the experimental parameters are provided in Table 5. All combinations of parameters listed were applied to the experimental setup at a cooling temperature of 240 °C. The experimental plan was executed in seven phases as follows:

1.) **Ramp-up at benchmark conditions at 50 bar:**
\[ p = 50 \text{ bar}; \ T_{\text{cool}} = 240 \degree \text{C}; \ \text{SN} = 4.0; \ \text{COR} = 0.9; \ \text{GHSV} = 12,000 \text{ h}^{-1} \]

2.) **Parameter variation at 50 bar and 240 °C:**
\[ p = 50 \text{ bar}; \ T_{\text{cool}} = 240 \degree \text{C}; 2.0 \leq \text{SN} \leq 8.0; \ \text{SN} \leq \text{COR} \leq 0.95; 5,000 \text{ h}^{-1} \leq \text{GHSV} \leq 12,000 \text{ h}^{-1} \]

3.) **Parameter variation at 65 bar and 240 °C:**
\[ p = 65 \text{ bar}; \ T_{\text{cool}} = 240 \degree \text{C}; 2.0 \leq \text{SN} \leq 8.0; \ \text{SN} \leq \text{COR} \leq 0.95; 6,000 \text{ h}^{-1} \leq \text{GHSV} \leq 12,000 \text{ h}^{-1} \]

4.) **Parameter variation at 80 bar and 240 °C:**
\[ p = 80 \text{ bar}; \ T_{\text{cool}} = 240 \degree \text{C}; 2.0 \leq \text{SN} \leq 8.0; \ \text{SN} \leq \text{COR} \leq 0.95; 6,000 \text{ h}^{-1} \leq \text{GHSV} \leq 12,000 \text{ h}^{-1} \]

5.) **Parameter variation at COR = 0.98 and 240 °C:**
\[ p = 50 \text{ bar} \leq p \leq 80 \text{ bar}; \ T_{\text{cool}} = 240 \degree \text{C}; 2.0 \leq \text{SN} \leq 8.0; \ \text{SN} \leq \text{COR} \leq 0.98; 6,000 \text{ h}^{-1} \leq \text{GHSV} \leq 12,000 \text{ h}^{-1} \]

6.) **Parameter variation at 50 bar and 220 °C:**
\[ p = 50 \text{ bar}; \ T_{\text{cool}} = 220 \degree \text{C}; 2.0 \leq \text{SN} \leq 8.0; \ \text{SN} \leq \text{COR} \leq 0.95; 6,000 \text{ h}^{-1} \leq \text{GHSV} \leq 12,000 \text{ h}^{-1} \]

7.) **Benchmark at conditions of phase (1)**

During phase 1.) the benchmark conditions were held constant for 56 h. In phase 5.) COR was held constant at 0.98 at the three pressure levels considered. This variation was not included into phases 2.) to 4.) as catalyst deactivation was expected during
these experiments due to the high CO₂ content. After the parameter variation was terminated, the benchmark point of phase 1) was held constant for another 12 h.

Validation and parameter fitting

In order to validate the behaviour of literature kinetic models in comparison against the data acquired within this study and to optimize their behaviour by a parameter variation, information on gas phase composition and temperature profile for each working point were fed into a validation library. A reactor simulation was carried out for all documented working points to determine deviations between model and experiment. A multi-criterial optimization changing the parameters of the utilized kinetic models, i.e. kinetic constants and adsorption constants, was executed in order to minimize the deviation between the reactor simulation and the experimental data. The objective function \( f(x) \) for the parameter fitting was formulated as the sum of the weighted root mean square errors for temperature profile (\( RMSE_{\text{Temp Profile}} \)), for hot spot temperature (\( RMSE_{\text{Ts}} \)) and gas phase composition at reactor outlet (\( RMSE_{\gamma} \)):

\[
f(x) = \alpha \cdot RMSE_{\text{Temp Profile}} + \beta \cdot RMSE_{\text{Ts}} + \gamma \cdot RMSE_{\gamma}
\]

(35)

The RMSE in Eq. (35) were calculated as follows:

\[
RMSE_{\text{Temp Profile}} = \sqrt{\frac{\sum_{i=1}^{N_{\text{data pt}}} (T_{\text{exp},i} - T_{\text{sim},i})^2}{N_{\text{data pt}}}}
\]

\[
RMSE_{\text{Ts}} = \sqrt{\frac{\sum_{i=1}^{N_{\text{data pt}}} (T_{\text{exp},i} - T_{\text{sim},i})^2}{N_{\text{data pt}}}}
\]

\[
RMSE_{\gamma} = \sqrt{\frac{\sum_{i=1}^{N_{\text{data pt}}} (y_{\text{exp},i} - y_{\text{sim},i})^2}{N_{\text{data pt}}}}
\]

The weighting factors \( \alpha, \beta \) and \( \gamma \) were fixed to constant values of \( \alpha = 2 \, K^{-1} \), \( \beta = 3 \, K^{-1} \) and \( \gamma = 25 \) by an empirical approach in order to balance the RMSE of temperature and composition to comparable numerical values.

Discussion of the impact on industrial reactor design

In order to demonstrate the impact of the new kinetic model on the industrial scale reactor in comparison to our previously published model, a simulation study was performed analysing hot spot position and temperature as well as product composition, in an industrial reactor simulation at the three pressure levels 50 bar, 65 bar and 80 bar at GHSV = 6,000 h⁻¹. For the gas composition the range applied in the experimental campaign was considered (see Table 5). Inlet and cooling temperatures were set to 240 °C.

Results and Discussion

Experimental results

Experimental data obtained from the miniplant setup indicated strong sensitivities of hot spot temperature, product composition and space time yield (STY) towards pressure, stoichiometry and COR. However, the measurement campaign was overlaid by a continuous deactivation of the catalyst. In Figure 4 STY is plotted over experimental time-on-stream (ToS) for the benchmark composition of \( SN = 4.0 \) and \( COR = 0.9 \) at the three pressure levels as well as the two cooling temperatures applied in this study. The graph indicates that STY stabilized during ramp up after approx. 50 h ToS. However, stronger deactivation of the catalyst was observed during the experimental phase at 80 bar (phase 4) and \( COR = 0.98 \) (phase 5). As both, the highest temperatures and the highest water contents were measured during these phases, based on these observations it can be concluded that the deactivation of the catalyst was mainly correlated to these two factors. This is in good agreement to the work of Fichtl et al. who considered hydrothermal degradation of the active sites as the main reason.
for catalyst deactivation in cleaned syngas. However, their group showed the necessity for longer experimental campaigns exceeding 1600 h ToS to obtain satisfactory information about deactivation kinetics. As this, though, was not in the scope of our study, the influence of catalyst deactivation was not yet included within our study consequently leading towards inaccuracies for the kinetic fitting. Future research is planned in order to derive advanced axially resolved deactivation kinetics using the experimental setup described herein.

In Figure 5 the molar fractions of water and methanol obtained from the experiments at GHSV = 12,000 h⁻¹ and a cooling temperature of 240 °C at the three pressure levels for COR = 0.7 (left) and COR = 0.95 (right) are depicted over SN. Thermodynamic equilibrium for the data points was calculated including Le Chatelier’s principle, increased synthesis due to higher reaction kinetics of rWGS. As expected, production did reach the thermodynamic equilibrium, probably inhibiting effect of high water partial pressures that was already recorded in literature.

Besides product concentration, another indicator for the catalytic activity can be gathered from the temperature profile inside the reactor. In Figure 6 the hot spot temperatures for the three pressure levels at GHSV = 12,000 h⁻¹ and COR = 0.7 and the particle distribution along the temperature sensor, a clear temperature difference between equilibrium and measured molar fraction of methanol, all experiments at this GHSV were carried out inside the kinetic regime of the methanol reaction. However, water production did reach the thermodynamic equilibrium, probably due to higher reaction kinetics of rWGS. As expected considering Le Chatelier’s principle, increased synthesis pressures led towards increased equilibrium molar fractions of methanol and water and consequently to higher reaction kinetics due to an enhanced driving force. The highest methanol molar fraction was obtained at COR = 0.7 and SN = 2.0. While at COR = 0.7 an increase of SN led towards a decrease of methanol molar fraction, the molar fraction of methanol was not sensitive to SN at COR = 0.95. This finding can be explained with the rate sensitivity of hot spot position towards SN could not be derived.

Figure 4: Trend of the space time yield over time-on-stream at benchmark conditions CON = 0.9, SN = 4.0; GHSV = 12,000 h⁻¹ at 50 bar, 65 bar and 80 bar at T_out = 240 °C and T_out = 220 °C. Sectors marked: 1) ramp up, benchmark at 50 bar, 240 °C; 2) experimental plan at 50 bar and 240 °C; 3) experimental plan at 65 bar and 240 °C; 4) experimental plan at 80 bar and 240 °C; 5) variation of SN at COR = 0.98, 240 °C and 50 bar to 80 bar; 6) experimental plan at 50 bar, 220 °C; 7) benchmark at 50 bar, 240 °C.

Figure 5: Equilibrium and measured molar fraction of methanol (black) and water (grey) at COR = 0.7 (left) and COR = 0.95 (right); Equilibrium molar fractions at 50 bar (dotted), 65 bar (dashed) and 80 bar (solid); measured molar fractions of methanol and water at GHSV = 12,000 h⁻¹ at 50 bar (triangle), 65 bar (diamond) and 80 bar (circle).

Figure 6: Hot spot temperature measured at COR = 0.7 (black) and COR = 0.95 (grey) and GHSV = 12,000 h⁻¹, pressure levels: 50 bar (triangle), 65 bar (diamond) and 80 bar (circle).
At COR = 0.7 an increase of SN led towards a decrease of hot spot temperature at 65 bar and 80 bar, whereas it was almost constant at 50 bar. This can be explained by chemical equilibrium of methanol synthesis decreasing by rising SN and temperature as well as the acceleration of reaction kinetics at increased temperature and pressure. Most probably hot spot temperature was limited by chemical equilibrium at 65 bar and 80 bar when SN exceeded a value of 3.0. As temperature downstream the hot spot approaches cooling temperature and therefore higher equilibrium conversions at simultaneously slower reaction kinetics, the difference between equilibrium and measured molar fractions in Figure 5 can be explained. While at COR = 0.7 an increase of synthesis pressure from 50 bar to 65 bar as well as from 65 bar to 80 bar increased the hot spot temperature over at least 7 K, at COR = 0.95 only a small rise of hot spot temperature for less than 3.5 K was measured. Moreover, sensitivity towards SN was weaker at COR = 0.95 with the highest hot spot temperature obtained at SN = 5.0 for all three pressure levels. Overall, the sensitivities of hot spot temperatures towards COR and SN are in good agreement to the results of our previous study, where the effect of feed gas composition towards temperature profile inside the reactor was discussed.27, 27

NMR side-product analysis of the liquid product showed the presence of low concentrations of ethanol, propanol and formic acid, which is in good agreement to Göhna et al. who analysed the product-compositions of CO₂-based methanol synthesis.71 However, as non-condensable side-products as methane and DME could not be trapped inside the liquid phase, no comprehensive analysis could be drawn from the liquid phase measurements executed. Further side-product gas phase measurements utilizing a FTIR with a longer optical path length to identify possible traces of these components could be applied in future studies.

Overall, the experimental results obtained from the miniplant setup were plausible regarding the trends in hot spot temperature and product composition. Therefore, the authors are confident that the measured data provides a reliable data basis for the validation and adjustment of kinetic models.

Validation of literature kinetics

In order to discuss the ability of literature kinetic models for a description of the measured data obtained from the miniplant setup, reactor simulations using the kinetic models as proposed by Graaf27, Bussche15 and in our previous study, hereon denoted Nestler27, were performed for all experimental working points. For the sake of clarity, the kinetic models with the parameter set applied as published in literature are hereon labelled with the index “original”. In Figure 7 parity plots for the three models are provided for the product molar fractions of methanol (A) and water (B) as well as hot spot temperature (C) and position (D), with a confidence interval of 10%. The graphs for the outlet molar fraction of water and methanol show a high level of agreement between experiment and simulation in terms of the kinetic model Nestleroriginal. The models Busscheoriginal and Graaforiginal, however, show strong deviations from the experiments with the tendency of underestimated reaction kinetics.

Interestingly, none of the models considered in Figure 7 was able to precisely describe the thermal behaviour of the reactor. Even though the hot spot temperatures of all kinetic models lie within the 10% confidence interval, position of the hot spot was estimated further downstream in the catalyst bed for all kinetic models considered here. In Table 6 the objective function obtained from Eq. (35) is shown for the three original models considered in this study together with the respective RMSE-values.

Table 6: Objective function and RMSEs of the original kinetic models over the experimental data points.

| Parameter | Unit | Graaforiginal | Busscheoriginal | Nestleroriginal |
|-----------|------|---------------|----------------|----------------|
| f(x)      | -    | 52.06         | 59.54          | 24.25          |
| RMSEprofile K | 2.3 | 2.5           | 2.6            |
| RMSEprof K | 9.2  | 8.4           | 4.0            |
| RMSE%     | %    | 0.79          | 1.18           | 0.27           |

The RMSE values of the product composition prove the high accuracy of the Nestleroriginal model for calculation of the product composition in comparison to the literature standards Graaforiginal and Busscheoriginal. While these models predict product composition with a mean error of 0.79% and 1.18%, respectively, a smaller mean error of 0.22% is obtained when the Nestleroriginal model is applied. However, hot spot temperature of this model is still predicted with a mean error of 4 K. As the temperature profile is coupled with the conversion of synthesis gas towards methanol, wrong outlet concentrations could be calculated when the kinetic models discussed in this section are transferred towards different
reactor geometries, working conditions or even other reactor types, e.g. an adiabatic quench bed reactor. Even though, the kinetic model previously published by our group delivers a satisfactory description of the outlet concentration for the experimental conditions applied, high deviations could be the case, especially when the kinetic model is used for high COR and higher GHSV. This finding emphasizes the necessity for a highly resolved axial measurement in experimental kinetic campaigns, analogous to the temperature profile along the reactor as presented in this work. Moreover, an accurate prediction of the temperature profile is necessary for reactor design, especially when syngas with higher CO contents leading to higher hot spot temperatures is fed to the reactor.

Fitted kinetic models

In order to enhance the applicability of the kinetic models described previously, their semi-empirical parameters (see Eq. (20) and (21)) were fitted to the experimental results measured at the miniplant. The parameter fitting was subjected to the weighting factors in Eq. (35). Other weighting factors could influence the fitting result along the Pareto front of the optimization problem.2 The kinetic models refitted to the experimental data will be denoted with the index “fit” hereafter. In Table 7 the results of the parameter fitting utilizing the previously defined weight factors are listed by means of the objective function and the respective RMSE values.

Table 7: Objective function and RMSEs of the fitted kinetic models over the experimental data points.

| Parameter | Unit | Graaffit | Busschefit | Nestlerfit |
|-----------|------|----------|------------|------------|
| f(x)      |      | 17.50    | 24.97      | 17.93      |
| RMSE_f(i,profile) | K | 1.4 | 1.8 | 1.5 |
| RMSE_f(T,hs) | K | 1.7 | 3.4 | 1.8 |
| RMSE_f(T) | % | 0.38 | 0.45 | 0.38 |

Comparison of the fitted kinetic models shows comparable remaining errors for the models Graaffit and Nestlerfit, while for the model Busschefit larger deviations remain for temperature profile and product concentration. This can be explained by the reaction mechanisms and rds of the kinetic models. Graaffit and Nestlerfit rely on a common mechanism and similar rds, however with Nestlerfit not considering CO-hydrogenation. In contrast, Bussche’s rate equation is based on a different mechanism. Due to the high remaining errors after the parameter fitting (compare Table 7) the rate equations of the Bussche-model were found not applicable for the description of methanol synthesis kinetics on the catalyst considered in this study.

The remaining RMSE values show that the fitted models Graaffit and Nestlerfit predict the temperature profile with a mean error of 1.4 K or 1.5 K, respectively, and therefore with a higher accuracy than the original literature models. A deeper look into the reaction velocities of the fitted kinetic models at \( T_{\text{cool}} = 240^\circ \text{C} \) over the whole considered parameter range showed, that CO-hydrogenation of the Graaf fit model can be neglected due to a very small reaction rate \( \left( \frac{r_{\text{CO2}}}{r_{\text{WGS}}} > 3.2 \times 10^{-3} \text{ mol s}^{-1} \text{kg}_{\text{cat}}^{-1} \right) \). Due to this finding, it can be stated, that CO-hydrogenation can be neglected for the description of the kinetic behaviour inside the reactor, which is in good agreement to the findings of the scientific community.11,12 Consequently, the kinetic model Nestlerfit will be used throughout the following discussion of this publication.

The set of fitted kinetic parameters for the proposed kinetic model based on the rate equations of Eq. (16) and Eq. (17) is given in Table 8. The parameter sets of the other kinetic models fitted to the experimental data are provided in the supplementary material.

Table 8: Parameters for the proposed kinetic model Nestlerfit.

| Parameter | Unit | Proposed kinetic parameters |
|-----------|------|-----------------------------|
| \( k_1 \) | mol kg\(^{-1}\)s\(^{-1}\)Pa\(^{-1}\) | 2.385 \( \times 10^{-5} \) \( \exp \left( \frac{-14.09}{R \cdot T} \right) \) |
| \( k_2 \) | mol kg\(^{-1}\)s\(^{-1}\)Pa\(^{0.5}\) | 244.433 \( \exp \left( \frac{-53.741}{R \cdot T} \right) \) |
| \( K_1 \) | Pa\(^{-1}\) | 1.440 \( \times 10^{-17} \) \( \exp \left( \frac{-570}{R \cdot T} \right) \) |
| \( K_2 \) | Pa\(^{-1}\) | 4.223 \( \times 10^{-6} \) |
| \( K_3 \) | Pa\(^{0.5}\) | 6.407 \( \times 10^{-13} \) \( \exp \left( \frac{126.843}{R \cdot T} \right) \) |

In Figure 8 the parity plots for the outlet concentrations of methanol (A), water (B) as well as the hot spot temperature (C) and position (D) simulated with the proposed model are provided. The graphs indicate that the description of both, hot spot position and temperature were improved significantly in comparison to the original model (compare Figure 7). However, while the description of the temperature profile was enhanced.
with the proposed model, a slightly higher error can be observed regarding the composition of the products methanol and water. This is most likely due to inaccuracies in the measurements of axial temperature profile and product composition. Besides, the remaining error could be a consequence of inaccuracies in the reactor model, e.g. the diffusion or heat transfer sub-models. Application of the validation methodology presented within this study to other reactor geometries could help identifying possible simulation issues and improve the simulation platform. Despite the slightly lower accuracy of the proposed model in comparison to Nestler\textsubscript{original} for the calculation of product composition, it is worth pointing out, that the correct description of reaction kinetics along the reactor is vital to enable a reliable transfer of the kinetic model towards industrial scale. To the best of the authors’ knowledge, the herein proposed kinetic model delivers such a description and is therefore of a high value for such reactor design problems. However, the validity of the herein proposed kinetic model was only confirmed within the parameter range applied for the experimental campaign (compare Table 5). Expansion of the validated parameter range should only be applied with caution\textsuperscript{27}. More experimental data will be obtained from the miniplant for a wider and industrially relevant validity range in future work.

Impact on industrial scale

To quantify the behaviour of the herein proposed kinetic model on the industrial scale, a comprehensive simulation study was executed. As our previously published kinetic model\textsuperscript{27} was based on a similar catalyst, though exclusively based on the measurement of the outlet concentration of a kinetic reactor\textsuperscript{27}, comparison to this model is capable of showing the impact of the herein proposed validation approach. In Figure 9 industrial reactor simulations applying both, our previous kinetic model Nestler\textsubscript{original} and the proposed adapted kinetic model Nestler\textsubscript{fit} are compared by means of hot spot temperature (A, B) and position (C, D) as well as methanol (E, F) and water outlet molar fraction (G, H) at synthesis pressures of 50 bar (left side) and 80 bar (right side). The graphs A and B indicate a lower sensitivity of the Nestler\textsubscript{fit} model with regard to the dependency of hot spot position and temperature towards COR in comparison to Nestler\textsubscript{original}. While hot spot temperatures of both models are comparable at COR = 0.8 the proposed kinetic model shows lower hot spot temperatures at COR = 0.7 and increased temperatures at higher COR. As expected from the comparison of the parity plots of Nestler\textsubscript{original} model and Nestler\textsubscript{fit} model in Figure 7 and Figure 8, respectively, high deviations between the kinetic models are observed with regard to hot spot position. This shows that large inaccuracies on the industrial reactor scale can be obtained with kinetic models derived from experimental data measured in traditional integral reactors. Differential measurement of concentration or, as presented here, highly resolved temperature measurements add information to the data set which are advantageous when a transfer from lab to industrial scale is performed. Looking at the product molar fraction of methanol (E, F) and water (G, H) increasing deviations between the two models are present with decreasing SN. This is probably due to larger deviations in hot spot position predicted with decreasing SN. On the one hand this finding again shows the importance of the interlink between a correct kinetic axial description and accurate calculation of product formation. On the other hand, detailed knowledge of the product composition at the reactor exit is of high importance, when the synthesis reactor is embedded in a loop process.

Optimal design for the miniplant setup

In order to optimize the miniplant geometry for an improved agreement between industrial and miniplant scale, scale down from industrial scale to the miniplant dimensions was repeated applying the Nestler\textsubscript{fit} kinetic model. In Figure 10 the optimized reactor diameters determined at GHSV = 9,000 h\textsuperscript{-1} and the pressure levels of 50 bar (A) and 80 bar (B) are shown in a 2D contour plot. The graphs indicate that scale down of the industrial reactor to miniplant scale is correlated to the working range applied. While pressure and COR reveal higher sensitivities towards optimal reactor dimensions, SN does affect the diameter less significantly. With regard to the methodology...
applied an inner reactor diameter of 9 mm ≤ d_in ≤ 12 mm would be beneficial for the miniplant setup to improve the similarity towards the industrial reactor scale. Moreover, the smaller reactor diameter would lead to a better heat removal from the reactor and consequently enable the setup to be used for syngas with lower COR. However, wall effects (Eq. (31)) as well as other relevant design criteria must be considered when the geometry of the miniplant reactor is changed to the dimension proposed here.

As the implementation of Thiele modulus for the description of the diffusion showed to significantly influence the results of the scale down, further research will be necessary in order to validate the diffusion model against experimental data. This could be done by introduction of larger catalyst particles into the miniplant reactor in future work.

**Capability of the miniplant for the dynamic reactor analysis**

Besides steady state validation, the miniplant setup introduced within this work clearly provides the opportunity to validate reaction kinetics under transient conditions, i.e. fluctuating gas quantity or composition. In Figure 11 the change in the reactor’s temperature profile during an exemplary load change at a pressure level of 80 bar and T cool = 240 °C from COR = 0.8; SN = 2.0; GHSV = 6,000 h⁻¹ towards COR = 0.9; SN = 4.0; GHSV = 12,000 h⁻¹ is shown. The graph indicates the displacement of the hot spot inside the reactor further upstream and the decrease of hot spot temperature as a consequence of the increased CO₂ content in the feed gas. Further studies are planned in order to validate the dynamic behaviour of the miniplant reactor by a dynamic reactor simulation.

**Conclusions**

In this study, a novel approach for kinetic model validation and parameter estimation using experimental data from a miniplant setup featuring a highly resolved fibre optic temperature profile in a polytropic miniplant in combination with FTIR product composition measurement was presented. The experimental data obtained from the miniplant reactor are highly correlated to an industrial scale reactor according to the simulation platform applied in this work. Comparison of the experimental data to the reactor simulation of the miniplant using different kinetic models from literature showed the validity of our previously published kinetic model²⁷ by means of the product gas composition. However, comparison of the temperature profiles obtained by reactor simulation towards the experimental data proved the necessity for highly resolved axial measurement to obtain a satisfactory kinetic description. A parameter fitting minimizing the deviation between the experimental data and the simulation was carried out for the rate equations proposed by Graaf¹³, Bussche¹⁵ and Nestler²⁷. Remaining discrepancies between the adapted model by Bussche and the experimental data proved that the rate equation proposed by the authors is not capable of describing the reaction kinetics of the catalyst analysed in this work. In contrast, the refitted models by Graaf and Nestler showed a similar quality for the description of the reaction kinetics. As the reaction rate of CO-hydrogenation of the refitted Graaf model was by orders of magnitude below that of CO₂-hydrogenation, it can be concluded that the combination of RWGS and CO₂-hydrogenation is sufficient for the kinetic description in the...
parameter range considered. Based on the kinetic rate equation formulated within our previous work\textsuperscript{27}, a new data set of kinetic parameters was fitted.

A sensitivity analysis performed in the valid parameter range of the herein proposed kinetic model proved the advantage of the herein proposed methodology over classic kinetic fixed bed measurements for scale-up to an industrial reactor. In order to obtain even higher comparability of the miniplant towards industrial scale, the diameter of the miniplant reactor could be adapted in future work based on the simulation-based scale down presented in this work utilizing the updated kinetic model proposed within this study.

Moreover, the data obtained from the miniplant setup was found highly promising for the analysis of a dynamically operated polytropic methanol synthesis reactor. Further work will be carried out to validate a dynamic reactor model against experimental data obtained under transient conditions. Key issues in our work arise from catalyst deactivation during the experimental study. Validation and adaption of deactivation models using the miniplant setup are important tasks to increase the accuracy of the herein proposed methodology in future work. Moreover, application of the methodology for the spatially resolved validation of a diffusion model will be examined.

Even though the proposed kinetic model shows a high level of agreement towards the experimental data, further research regarding a more appropriate mechanistic description of methanol synthesis could be helpful to deliver an even better description of the reaction kinetics. The authors are confident that the herein applied experimental setup will be a helpful tool in order to clarify the mechanistic nature of methanol synthesis in future work.

To the best of the authors’ knowledge the herein proposed novel approach for the validation of reaction kinetics of fixed bed reactions is a significant improvement as it offers an enhanced methodology for bridging between experimental and industrial reactors. Further studies could be carried out transferring this methodology towards other fixed bed syntheses. Moreover, the consideration of radial effects by a two-dimensional simulation could positively affect the quality of the kinetic model obtained and should therefore be investigated in subsequent work.

Nomenclature

| Symbol | Definition |
|--------|------------|
| D_{ax} | Axial diffusion coefficient $[m^2 s^{-1}]$ |
| D_{ij} | Binary diffusion coefficient of component i and j $[m^2 s^{-1}]$ |
| D_{K,i} | Knudsen diffusion coefficient of component i $[m^2 s^{-1}]$ |
| D_{em,i} | Effective diffusion coefficient of component i $[m^2 s^{-1}]$ |
| EoS | Equation of State |
| GHSV | Gas hourly space velocity $[h^{-1}]$ |
| h_{cat} | Height of catalyst bed $[m]$ |
| ΔH_f | Enthalpy of formation under standard condition $[kJ mol^{-1}]$ |
| k_i | Reaction rate constant of reaction i $[mol s^{-1} m^{-3} Pa^{-1}]$ |
| K_i | Adsorption constant of component i $[variable unit]$ |
| K_{eq,i} | Equilibrium constant of reaction i $[variable unit]$ |
| K_{equ,i} | Pseudo equilibrium constant $[-]$ |
| n | Molar flow $[mol s^{-1}]$ |
| N | Number $[-]$ |
| p | Pressure $[Pa]$ |
| r_{eff,i} | Particle reaction velocity of reaction i $[mol kg^{-1} s^{-1}]$ |
| r_i | Intrinsic reaction velocity of reaction i $[mol kg^{-1} s^{-1}]$ |
| R | Universal gas constant $[J mol^{-1} K^{-1}]$ |
| rds | Rate determining step |
| Re_p | Particle Reynolds number $[-]$ |
| RMSE | Root mean square error |
| rWGS | Reverse water-gas-shift reaction |
| SN | Stoichiometric number $[-]$ |
| SRK | Soave-Redlich-Kwong |
| STY | Space time yield $[kg_{MeOH} L_{cat}^{-1} s^{-1}]$ |
| t | Time $[s]$ |
| T | Temperature $[K]$ |
| u_0 | Empty tube gas velocity $[m s^{-1}]$ |
| U | Heat transfer coefficient $[W m^{-2} K^{-1}]$ |
| V | Volumetric flow rate $[m^3 h^{-1}]$ |
| V | Volume $[m^3]$ |
| WGS | Water-gas-shift reaction |
| x | Axial length $[m]$ |
| y_i | Molar fraction of component $[-]$ |

Greek letters

| Symbol | Definition |
|--------|------------|
| α | Weight factor of temperature profile $[K^{-1}]$ |
| β | Weight factor of hot spot $[K^{-1}]$ |
| γ | Weight factor of molar fraction $[mol fraction]$ |
| ε | Porosity $[-]$ |
| η_{eff,i} | Efficiency factor of reaction i $[-]$ |
| λ | Heat conduction coefficient $[W m^{-1} K^{-1}]$ |
| u_j | Stoichiometric factor of component j $[-]$ |
| p | Density $[kg m^{-3}]$ |
| τ | Tortuosity of the catalyst particle $[-]$ |
| ν | Kinematic viscosity $[m^2 s^{-1}]$ |
| φ | Reactor-particle diameter ratio $[-]$ |
| Φ_{M,i} | Thiele modulus $[-]$ |

Indices

| Symbol | Definition |
|--------|------------|
| bulk | Bulk phase |
| cat | Catalyst in the reactor |
ARTICLE

conf comp cool data pt exp ext f (g) H2 H2O hs inc ind int MeOH miniplant norm p profile rad sim wall T tot x y

Component Cooling medium Data points Experimental External Fluid phase Gas phase Hydrogen Water Hot spot Increment Industrial scale Internal Miniplant scale Norm conditions Particle Profile Radial Simulated Wall Temperature Total Axial dimension Molar fraction

Conflicts of interest
There are no conflicts to declare.

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