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Supporting Information available online

1 Introduction

The growing energy demand of the developed and emerging economies is still mainly satisfied by oil, gas, and coal. However, the consequences of the climate change as a result of fossil fuel consumption, i.e., high CO_2 emissions, are becoming more and more obvious. Thus, renewable energy sources such as wind and solar enabling the substitution of fossil fuels are gaining importance, but their drawback is still the gap in demand and supply. In times of low winds and low or no sun radiation, electricity production lacks the demand, whereas on the other hand windy and sunny periods may yield a surplus of available electrical energy. Consequently, energy storage systems are needed to level the fluctuating production and use of energy [1–3].

One option for storage of surplus electricity is the conversion into chemical energy by the power-to-gas (PtG) technology [4]. In a first step, renewable H_2 is produced via electrolysis. However, H_2 requires special materials for storage tanks and the volumetric energy density of H_2 is low, e.g., compared to methane [5]. Here, subsequent conversion of H_2 with CO₂ into methane (synthetic natural gas, SNG) comes into play:

$$\mathrm{CO}_2 + 4 \mathrm{H}_2 \rightleftharpoons \mathrm{CH}_4 + 2 \mathrm{H}_2 \mathrm{O} \quad \Delta H_\mathrm{R} = -165 \mathrm{kJ} \mathrm{mol}^{-1} \quad (1)$$

SNG production has two advantages: CO_2 can be utilized and a technically mature storage and transport infrastructure for natural gas is already present in many countries [6]. SNG can substitute natural gas in existing applications, e.g., heat and power production or generation of syngas [7]. Hence, PtG provides an attractive link between the electrical and natural gas grid.

Beside CO_2 from flue gases of power plants or of the steel and cement industry [6], biogas is also an interesting CO_2 source. It contains about 60 % CH₄ and 40 % CO₂ and could be directly upgraded into SNG by addition of the respective amount of hydrogen (according to Eq. (1) 4 mol H₂ per mol CO_2) and subsequent methanation [6, 8, 9].

The fluctuating production of renewable H_2 may lead to the need of temporary H_2 (and CO₂) storage facilities in a PtG plant. In addition, the PtG chain may also require novel concepts for reactor operation which are optimized for smaller plant sizes and for intermittent or dynamic operation of the methanation unit. If the reactor is, e.g., suitable for a variation of the volume rate of the feed gas, this would help to reduce the size of gas storage tanks [10].

Several phenomena are reported for reactors in dynamic operation, which make stable and safe operation as in case of the exothermic methanation difficult: Experimental data and numerical modeling show wrong-way behavior, travelling hotspots and variations in the conversion of the syngas in cooled fixed-bed reactors [11]. Further simulations indicate also advantages of dynamic reactor operation such as a

Transient Effects during Dynamic Operation of a Wall-Cooled Fixed-Bed Reactor for CO₂ Methanation

The power-to-gas process is an option to transform fluctuating renewable electric energy into methane via water electrolysis and subsequent conversion of H_2 by methanation with CO_2 . The dynamic behavior of the methanation reactor may then be a critical aspect. The kinetics of CO_2 methanation on a Ni-catalyst were determined under isothermal and stationary conditions. Transient isothermal kinetic experiments showed a fast response of the rate on step changes of the concentrations of H_2 , CO_2 ; in case of H_2O , the response was delayed. Non-isothermal experiments were conducted in a wall-cooled fixed-bed reactor. Temperature profiles were measured and the effect of a changing volumetric flow was studied. The experimental data were compared with simulations by a transient reactor model.

Keywords: Dynamic modeling, Fixed-bed reactor, Methanation, Power-to-gas, Transient kinetics

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decreasing hotspot temperature [12] and higher conversion [13]. In literature optimizations for start-up, load changes, or mass and heat recycle are described [14–17], but steady-state kinetics are frequently assumed [12–17], although transient kinetic effects on the catalyst surface are also reported [18–21]. Hence, a reliable transient model that describes the main effects in dynamic operation of a reactor needs an experimental validation with regard to transient effects on the catalyst surface.

In this work, the kinetics of methanation were at first determined at isothermal steady-state conditions. With regard to transient operation it is necessary to know if steady-state kinetics can be assumed as transient kinetics, i.e., if the response of the reaction rate on changes of gas composition or temperature is fast. Therefore, step changes of the inlet gas concentrations of H_2 , CO_2 , and H_2O were performed to investigate the transient behavior of the catalyst. Moreover, dynamic and non-isothermal experiments with regard to a step change of the volume rate of the syngas were conducted in a wallcooled single-tube fixed-bed reactor. The experimental data were then compared with a suitable transient reactor model.

2 Experimental Setup and Procedure

2.1 Stationary and Transient Isothermal Kinetic Measurements

The setup for the isothermal kinetic measurements (Fig. S1 in Supporting Information) consists of a gas dosing unit, a small fixed-bed reactor, and a gas analysis. The feed rates of the gases H_2 , CO_2 , N_2 , and CH_4 were adjusted by mass flow controllers. Steam was added by a water saturator and the respective pipes were heated to prevent condensation. All experiments were conducted with a volumetric flow rate of $50 \text{ L} \text{ h}^{-1}$ (STP) and at 1 bar.

The lab-scale reactor was externally heated and cooled by a thermo-oil thermostat to provide isothermal conditions. The reactor had a diameter of only 1.4 cm and was charged with 1.5 g (1.8 g for transient experiments) of catalyst particles (NiSat 310 RS from Clariant) diluted with 7.5 g of inert quartz sand to ensure isothermal conditions. The bed length was about 5 cm. Data of the catalyst are given in Tab. 1.

For temperature measurements to prove isothermal conditions, a guiding tube for a thermocouple with a diameter of 2 mm was placed in the center of the tubular reactor.

Table 1. Characteristic data of the catalyst.

Parameter	Value
Particle diameter $d_{\rm p}$ [mm]	2.1
Particle density $ ho_{ m p}$ [kg m ⁻³]	3690
BET surface area [m ² g ⁻¹]	184
Average pore diameter [nm]	6.6
Ni content on Al ₂ SiO ₅ /Al ₂ O ₃ [wt %]	56.7
Heat capacity at 190 °C $c_{\rm p}~[{\rm J~kg^{-1}K^{-1}}]$	842

For the steady-state and isothermal kinetic experiments, the product gas passed through cooling traps at 0 °C and at -78 °C, respectively, and was then analyzed by a gas analyzer (Fisher-Rosemount MLT 2) for CO, CO₂, CH₄, and H₂. The experiments were conducted in a temperature range from 170 °C to 230 °C with syngas consisting of 4–20 % CO₂, 5–80 % H₂, 0–45 % H₂O, and 0–25 % CH₄. Nitrogen was only added to adjust the concentrations, e.g., to vary the concentration of only one reactant.

For the transient kinetic experiments the product gas was analyzed by a mass spectrometer (Pfeiffer MS Thermostar). By this, the response time of the gas analysis to a change of the adjustment of the mass flow controllers was lower than 10 s. In addition, a small amount of helium $(1 L (STP) h^{-1}; typically 2\%)$ of the total gas flow) was continuously added as an inert internal standard allowing the detection of volume flow fluctuations, e.g., an increasing signal of helium indicates a lower volume flow. Furthermore, the helium signal was used for the dead-time correction. The step change experiments between two concentrations were performed at 220 °C and the response signal was observed.

2.2 Transient Volume Flow Step Change Experiments with a Cooled Single-Tube Methanation Reactor

For the cooled single-tube methanation reactor (Fig. S2), a mass flow controller for a premixed syngas $(63 \% H_2, 15.3 \% CO_2, and 21.7 \% N_2)$ was applied. The reactor was a jacketed stainless-steel tube (inner diameter 2 cm, length 1 m). It was heated and cooled by an oil thermostat (LAUDA USH 400) and contains a 10-cm filling of glass spheres for consistent conditions at the inlet and subsequent 88-cm filling of catalyst. A guiding tube in the center of the reactor (3 mm diameter) over the total length was used to measure the axial temperature profile by means of a movable thermocouple.

Two cooling traps were placed downstream of the reactor to condense the product water, one at room temperature and the other one at 0 °C. The dry gas composition was analyzed by an infrared gas analyzer for CO, CO₂, as well as CH_4 and a thermal conductivity detector for H₂.

For axial temperature profile measurements, a sled driven by a stepper motor moved the thermocouple downwards the reactor. A computer controlled the mass flow controller in the same way as the stepper motor moved the thermocouple and recorded the measured temperature and gas concentration data from the gas analyzer. The experiment started with a volume flow of 60 Lh^{-1} for 45 min at a cooling temperature of 183 °C. Then, a step change of the flow to 7.5 Lh^{-1} was conducted for 45 min. This procedure was also done with a reverse step change ($7.5 \text{ Lh}^{-1} \rightarrow 60 \text{ Lh}^{-1}$).

To record the transient temperature profile with only one thermocouple (Fig. S3), both experiments were repeated 29 times at different positions of the thermocouple inside the reactor. Hence, the total experimental time of both step change experiments was 45 h. Within the first 10 cm of the reactor – the hotspot region – the temperature was measured in intervals of 1 cm. For the next 10 cm of the reactor length, the thermo-

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couple was moved in intervals of 2 cm and from this part of the reactor onwards to the end of the catalyst bed in the reactor in intervals of 5 cm.

3 Modeling a Wall-Cooled Fixed-Bed Methanation Reactor

An ideal one-dimensional plug-flow reactor was chosen as reactor model. This model neglects effects such as radial and axial dispersion of mass, radial velocity profiles, as well as radial temperature and concentration gradients within the fixed bed. The pressure drop over the fixed-bed reactor could be regarded as insignificant. The mass balance for each component i ($i = H_2$, CO₂, CH₄, H₂O, N₂) is then represented by:

$$\varepsilon \frac{\partial c_{\rm i}}{\partial t} = -\left(\frac{u_{\rm s} \partial c_{\rm i}}{\partial x} + \frac{c_{\rm i} \partial u_{\rm s}}{\partial x}\right) + \nu_{\rm i} \rho_{\rm bed} r_{\rm m} \tag{2}$$

The change in gas velocity derives from the non-stoichiometric reaction of CO₂ with H₂ and is a function of the reaction rate r_m (in mol CO₂ kg⁻¹s⁻¹) and the stoichiometric coefficients (Eq. (3)). Velocity changes due to the change of the axial temperature are small and were neglected.

$$\frac{\partial u_{\rm s}}{\partial x} = -2 \frac{r_{\rm m} \rho_{\rm bed}}{c_{\rm tot}} \tag{3}$$

For the energy balance, a pseudo-homogeneous model was used:

$$\left(\varepsilon \rho_{\rm g} c_{\rm p,g} + \rho_{\rm bed} c_{\rm p,bed} \right) \frac{\partial T}{\partial t} = -\rho_{\rm g} c_{\rm p,g} \left(\frac{(u_{\rm s} \partial T)}{\partial x} + \frac{T \partial u_{\rm s}}{\partial x} \right)$$

$$+ \frac{4}{d_{\rm r}} U_{\rm bed} (T_{\rm cool} - T) + \lambda_{\rm ax} \frac{\partial^2 T}{\partial x^2} - \Delta H_{\rm R} \rho_{\rm bed} r_{m}$$

$$(4)$$

The gas density $\rho_{\rm g}^{(1)}$ under reaction conditions was calculated using the reactor cooling temperature as reference. Both the thermal conductivity $\lambda_{\rm g}$ and viscosity $\eta_{\rm g}$ of the gas were calculated according to the Wassiljewa relation and the Mason Saxena modification [22]. The heat capacity $c_{\rm p}$ of the gas was determined using a correlation given in [23]. The stagnant thermal conductivity of the fixed bed $\lambda_{\rm bed}$, calculated on the basis of the model of Zehner-Bauer-Schlünder, was 0.27 W m⁻¹K⁻¹ [23]. Correlations for *Nu*, $\lambda_{\rm rad}$, and $\alpha_{\rm w}$ were also taken from [23].

In order to determine the radial heat conduction λ_{rad} and the heat transfer from the fixed-bed to the cooled reactor wall α_{w} , an overall heat transfer coefficient U_{bed} was defined [24]. The (radial) maximum temperature in the center of the reactor (measured in the experiments) was calculated based on the modeled mean temperature, and the values of λ_{rad} and α_w [24]. For the reaction rate r_m , the kinetic parameters determined at steady-state conditions (see Sect. 4.1) were used. Hence, the reactor model only accounts for transient thermal effects, while, e.g., transient kinetic effects like ad- and desorption kinetics are not considered. Mass transport limitations by pore 2403

diffusion were negligible, as proven by an estimation of the Thiele modulus. Values of the model parameters under the conditions of the experiments with the wall-cooled methanation reactor are depicted in Tab. 2.

 Table 2. Parameters used for the modeling of the wall-cooled fixed-bed methanation reactor.

Parameter	Value	
Reactor diameter $d_{\rm r}$ [cm]/length $L_{\rm r}$ [cm]	2/88	
Bulk porosity (ε)	0.82	
Stagnant thermal conductivity $(\lambda_{\rm bed})$ $[Wm^{-1}K^{-1}]$	0.27	
Total pressure (<i>p</i>) [bar]	1	
Cooling temperature (T_{cool}) [°C]	183	
Volume rate of syngas (STP) $[L h^{-1}]$	7.5	60
Nu (internal wall, depends on $\lambda_{\rm g}$ and thus on conversion) [–]	3.6-11.9	3.9-7.6
Radial thermal conductivity (λ_{rad}) [W m ⁻¹ K ⁻¹]	0.27	0.29
Heat transfer coefficient (α_W) [W m ⁻² K ⁻¹]	238	260
Overall heat transfer coefficient (U_{bed}) [W m ⁻² K ⁻¹]	74	80

4 Results and Discussion

4.1 Steady-State Kinetics of Methanation

Various approaches for the kinetics of CO_2 methanation are given in the literature [25–27]. In this work, a Langmuir-Hinshelwood-Hougen-Watson (LHHW) approach was chosen as it allowed the best prediction of the reaction rate over a wide range of conditions. The rate equation is taken from Koschanýs approach for a Ni/AlO_(x) catalyst [25]:

$$r_{\rm m,CO_2} = \frac{k_{\rm m} p_{\rm CO_2}^{0.5} p_{\rm H_2}^{0.5} \left(1 - \frac{p_{\rm CH_4} p_{\rm H_2} o^2}{p_{\rm CO_2} p_{\rm H_2^4} K_{p,Pa}}\right)}{\left(1 + K_1 p_{\rm CO_2}^{0.5} + K_2 p_{\rm H_2}^{0.5} + K_3 \frac{p_{\rm H_2} o}{p_{\rm H_2}^{0.5}}\right)^2}$$
(5)

The constants $k_{\rm m}$ and K_{1-3} depend on temperature and can be described by the Arrhenius (Eq. (6)) and van't Hoff equation (Eq. (7)), respectively.

$$k_{\rm m} = k_{0,\rm m} e^{\left(-\frac{{\rm E}_{\rm A}}{{\rm RT}}\right)} \tag{6}$$

$$K_{1-3} = k_{0,1-3}e^{\left(-\frac{\Delta_{\text{ads},1-3}H}{RT}\right)}$$
 (7)

The equilibrium constant K_p (in Pa⁻²) was calculated according to the equation developed by Aparicio [28] (with *T* in K):

¹⁾ List of symbols at the end of the paper.

$$K_{\rm p,Pa} = 137 \times 10^{-10} T^{-3.998} e^{\left(\frac{158.7 \times 10^3 \frac{J}{\rm mol}}{\rm RT}\right)}$$
(8)

Due to low conversion of CO_2 (<15%), differential conditions are assumed. A high conversion of CO_2 and thus a high concentration of steam were simulated by addition of H₂O to the feed gas. All experimentally determined parameters are listed in Tab. 3.

Table 3. Kinetic data of reaction rate constant and the three adsorption constants (see Eqs. (5)–(7)).

<i>k</i> _{0,m}	$9.2 \times 10^{-2} \operatorname{mol}(\text{kg s Pa})^{-1}$
$E_{\rm A}$	46.8 kJ mol ⁻¹
K _{0,1}	$4.5 \times 10^{-6} Pa^{-0.5}$
$\Delta_{\mathrm{ads},1}H$	-29.4 kJ mol ⁻¹
K _{0,2}	$1.7 \times 10^{-5} \mathrm{Pa}^{-0.5}$
$\Delta_{\mathrm{ads},2}H$	$-23.4 \text{ kJ mol}^{-1}$
K _{0,3}	$1.5 \times 10^{-6} \text{Pa}^{-0.5}$
$\Delta_{\rm ads,3}H$	-34.7 kJ mol ⁻¹

LHHW approaches contain different mechanistic assumptions, but there is still no agreement about the reaction steps actually occurring [29]. The reaction orders of H_2 and CO_2 (both 0.5) were determined by experiments with varied concentration of both components (for CO_2 see also Fig. 1). The order for CO_2 indicates a dissociative adsorption of CO_2 into CO and atomic O on the surface followed by CO methanation [18, 30]; the reaction order of H_2 can be explained by dissociative adsorption. Thus, CO_2 and H_2 each occupy two active sites. For SNG production, a high conversion of CO_2 is demanded. This yields significant amounts of CH_4 and H_2O , and product inhibition may occur. Thus, the influence of both product compounds on the methanation kinetics was also investigated. The experimental data gave no evidence for an influence of methane, but steam (water) is an inhibiting agent (Fig. 1).

Fig. 1 shows a decreasing reaction rate for constant H_2 and CO_2 partial pressures caused by an increasing partial pressure of water; this was also observed by Marwood et al. [19], who reported a decreasing CO coverage on the catalyst when water was added, and thus a reduction of adsorbed C-species. In the literature [25, 31, 32], the reaction order of water in the respective adsorption term is assumed to be unity. Table 4. With Eq. gas with Adsorpt $K_1p_{CO_2}^{0.5}$

Inhibition effects are represented by the adsorption terms for CO_2 , H_2 , and H_2O in the denominator of the LHHW approach. The strength of each inhibition is indicated in Tab. 4 by the comparison of the respective adsorption terms at a typical reaction temperature of 220 °C and different degrees of CO_2 conversion. Inhibition by H_2 and CO_2 declines with increasing conversion; for steam this is reverse.

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Figure 1. Influence of water vapor (diamonds) and CO₂ (circles) partial pressure on the reaction rate at 221 °C and 1 bar. H₂O variation: 10 % CO₂, 40 % H₂, 0–45 % H₂O in N₂; CO₂ variation: 4–20 % CO₂, 40 % H₂, 0 % H₂O in N₂; modified residence time: 60 s kg m⁻³ (221 °C, 1 bar).

The inhibiting effect of steam on the rate and conversion of CO₂, respectively, is also demonstrated in Fig. 2; just for comparison, the hypothetical case without H₂O inhibition is also depicted. Up to 20 % CO₂ conversion, the formed steam has almost no influence on the rate, but at higher conversion, inhibition becomes more and more relevant. Note that for the conditions given in Fig. 2 (220 °C, 1 bar, 20 % CO₂, 80 % H₂), the thermodynamic equilibrium conversion of CO₂ is 98.4 %. Almost full conversion can only be reached for lower temperatures (e.g., $X_{CO2,eq} = 99.3$ % at 180 °C), and higher temperatures are unfavorable with regard to CO₂ methanation (e.g., $X_{CO2,eq} = 94.9$ % for 300 °C).

Finally, a parity plot of the experimental results is presented in Fig. 3. The agreement of measured and calculated values is mainly in a range of ± 10 %.

Table 4. Comparison of the adsorption term and the reaction rate, calculated with Eq. (5)) at different conversion degrees of CO_2 at 220 °C, 1 bar, and a feed gas with 20 % CO_2 and 80 % H_2 .

Adsorption term / reaction rate	$X_{\rm CO2} = 25 \%$	$X_{\rm CO2} = 50 \%$	$X_{\rm CO2} = 75 \%$
$K_1 p_{\rm CO_2}^{0.5}$	0.75	0.65	0.49
$K_2 p_{\rm H_2}^{0.5}$	1.32	1.14	0.87
$K_3 \frac{p_{\rm H_2O}}{p_{\rm H_2}^{0.5}}$	0.31	0.79	1.80
$\left(1 + K_1 p_{\rm CO_2}^{0.5} + K_2 p_{\rm H_2}^{0.5} + K_3 \frac{p_{\rm H_2O}}{p_{\rm H_2}^{0.5}}\right)^2$	11.4	12.9	17.3
$r_{\rm m,CO2} [{\rm mol}{\rm kg_{cat}}^{-1}{\rm s}^{-1}]$	3.0×10^{-3}	2.0×10^{-3}	0.84×10^{-3}



Figure 2. Conversion of CO₂ as a function of the modified residence time with and for comparison without water inhibition term (K_3). CO₂ conversion and H₂O production were calculated by Eq. (5). Initial gas composition: 20 % CO₂, 80 % H₂, 0 % H₂O; 1 bar, $X_{CO2,eq} = 98.4$ %.



Figure 3. Parity plot $(170-230 \degree C, 1 \text{ bar}, \text{ modified residence time: } 60 \text{ s kg m}^{-3} \text{ at } 220 \degree C \text{ and } 1 \text{ bar}).$

4.2 Transient Response on the Kinetics on Partial Pressure Step Changes

In Sect. 4.1 the intrinsic steady-state kinetics were discussed. However, for dynamic operation it is necessary to know how fast the catalyst responds to changes of the reaction conditions such as the inlet gas composition, and whether and how fast steady-state conditions are again reached. Therefore, step change experiments were conducted. For example, if the time delay is very short, the kinetics determined at steady-state kinetics can be assumed to be valid also for a transient operation of a reactor, which simplifies the respective modeling.

The transient kinetic experiments were conducted under isothermal conditions. A mass spectrometer was used to analyze the feed and product gas. With helium as internal standard, changes of the volume flow during the initial phase of the step change and the dead time, respectively, could be identified and separated from real catalyst effects. The focus of the transient experiments was to investigate the influence of a stoichiometric change of the partial pressures of H_2 and CO_2 (Figs. 4 and 5) as well as of H_2O (Fig. 6).

Figure 4 reveals that the catalyst responds fast to a partial pressure step change of the feed gases CO_2 and H_2 from initially low to high values. The H_2 -to- CO_2 ratio was thereby kept constant at a value of 4. While the contents of CO_2 , H_2 , and CH_4 are already constant after about 25 s, the water content exhibits a certain delay before a stable value is achieved. Obviously, water, which is formed to a higher extent after the step change compared to before, adsorbs on the surface until the corresponding new adsorption equilibrium is reached. This corresponds to an initially slightly higher conversion (lower content) of CO_2 compared to the finally reached stationary value as water adsorption inhibits the reaction rate.



Figure 4. Molar content of CH₄, H₂O, and CO₂ before and after a partial pressure change of H₂ and CO₂ from low to high values at 221 °C and 1 bar. Dead time (<10 s) was corrected via helium standard (helium content: 2%). Gray area: adjustment of mass flow controllers (data points deleted). Residence time: 72 s kg m⁻³.

This behavior is similar but now in the reverse direction, if the step change is done from high to low partial pressures of CO_2 and H_2 (Fig. 5). Again, the new adsorption equilibrium for water is established with a certain delay as compared to H_2 and CO_2 , which now leads to an initially slightly lower conversion of CO_2 (more pronounced visible by lower methane content). Now, the amount of water formed after the step change is lower compared to before the change, and the inhibition by a water adsorbed on the surface, until the new adsorption equilibrium is established by desorption after about 1 min, leads to a decrease of the reaction rate.

So, ad- and desorption of steam is obviously the main effect prolonging the time required to reach steady-state conditions. In order to study this more explicit, only the steam content was now changed stepwise from 20% to a feed gas free of steam (Fig. 6). As expected, the content, i.e., the production rate, of



Figure 5. Molar content of CH₄, H₂O, and CO₂ before and after a partial pressure change of H₂ and CO from high to low values at 221 °C and 1 bar. Dead time (< 10 s) was corrected via helium standard (helium content: 2 %). Gray area: adjustment of mass flow controllers (data points deleted). Residence time: 72 s kg m^{-3} .



Figure 6. Molar content of methane and steam (in product gas) during a step change experiment with an initially high content of steam (20%) in the feed gas to a feed gas free of steam (221 °C, 1 bar). Gray area: adjustment of mass flow controllers, data points deleted. Steady-state conversion X_{CO2} : 6.8% (before step change), 8.8% (after step change). Modified residence time: 72 s kg m⁻³. Dead time (< 10 s) was corrected via helium standard (2% He).

methane is lower before the step change due to the higher (surface) concentration of water and the corresponding inhibiting effect by steam adsorption. After the step change from a rather high steam content of 20 % to a feed gas free of steam, methane production increases until after about 7 min the expected values (and accurately calculated by the kinetic model) for steady state are reached. The influence of the "delayed" steam ad- and desorption is also important to a certain extent for the transient operation of a wall-cooled fixed-bed reactor, as inspected is in the next Sect. 4.3.

4.3 Effects of Changing Volume Flow in a Wall-Cooled Fixed-Bed Reactor

Subsequent to the studies on the isothermal steady-state kinetics and on transient effects on the kinetics, experiments were conducted in a non-isothermal wall-cooled fixed-bed reactor. The transient behavior of the reactor was studied by a step change of the total volume rate. The reactor was also modeled as described in Sect. 3. For calculation of the conversion and heat production, the steady-state kinetics (see Sect. 4.1) were used. The experiment focused on biogas upgrading.

Biogas typically contains 60 % CH₄ and 40 % CO₂. Hence, at least 4 mol H₂ has to be admixed per mol CO₂ to achieve 100 % conversion of CO₂ to CH₄. Here, a slightly higher ratio of 4.2 was applied for adjusting a small H₂ excess to ensure full conversion. Methane was not added to the feed gas; instead, the respective amount of N₂ was added. Before presenting the dynamic measurements, the steady-state temperature profiles are illustrated in Fig. 7. The model describes the measured temperature profile and the location of the hotspot temperature satisfactorily. For the shorter residence time (higher volume rate), the model predicts a slightly higher CO₂ conversion, i.e., 69 % compared to 60 % measured, which may be explained by inaccuracies at high H₂O-to-H₂ ratios.



Figure 7. Steady-state axial temperature profile in a wall-cooled fixed-bed reactor; cooling/wall temperature (T_{cool}): 183 °C, pressure: 1 bar; inlet gas: 63 % H₂, 15.3 % CO₂, 21.7 % N₂; volume flow: 60/7.5 L h⁻¹ (STP) (diamond/triangle). Modified residence time: 6135/49079 s kg m⁻³ (220 °C, 1 bar).

The transient experiments with the wall-cooled fixed-bed reactor were of interest as the hotspot behavior during a load change is important for safe operation. In order to achieve the strongest effect, a step change was chosen, while in industrial practice the load change would be realized by a ramp. Note that the equilibrium conversion of CO^2 to methane is 100 % for the given low reaction temperature of around 190 °C. The transient temperature profiles were measured as described in Sect. 2.2.

In Fig. 8, the temperature profile at different times after the step change and in Fig. 9 only the measured and modeled hot-



Figure 8. Measured axial temperature profile of a wall-cooled fixed-bed reactor after a volume flow step change from 7.5 to $60 \text{ L} \text{ h}^{-1}$ (STP); inlet gas: $63 \% \text{ H}_2$, $15.3 \% \text{ CO}_2$, $21.7 \% \text{ N}_2$. P = 1 bar. Modified residence time: 6135 s kg m^{-3} (220 °C, 1 bar). Lines are guidance for the eye. Filled marks at hotspot position.



Figure 9. Hotspot temperature of a wall-cooled fixed-bed reactor after a volume flow step change from 7.5 to $60 \text{ L} \text{ h}^{-1}$ (STP); inlet gas: 63 % H₂, 15.3 % CO₂, 21.7 % N₂. Hotspot position at 7.5 L h⁻¹: 1.3 cm (model), 4 cm (measurement) with a steady-state conversion of 100 % (model and measurement). Modified residence time: 49079 s kg m⁻³ (220 °C, 1 bar). Hotspot at $60 \text{ L} \text{ h}^{-1}$: 2.4 cm (model), 5 cm (measurement) with a steady-state conversion of 60 % (see Fig. 7). Residence time: 6135 s kg m^{-3} (220 °C, 1 bar). Subfigure shows the modeled amount of water at both feed rates at different axial positions in the front part of the reactor at steady state.

spot temperatures (i.e., maximum axial temperatures as depicted as filled marks in Fig. 8) are displayed.

The model predicts a faster adjustment of the steady-state hotspot temperature after about 2 min, whereas 5 min are needed experimentally (Fig. 9). Furthermore, the model fails to describe the change of the hotspot in this interval. The model shows an immediate increase of the hotspot temperature, whereas in the experiment the temperature of the hotspot decreases initially and then rises subsequently. This is observed over the whole length of the reactor (see Fig. 8). The measured temperature rise at the rear end of the reactor at t = 60 s is probably due to the exothermic adsorption of CO₂ and H₂, which is not present before in the rear section prior to the step change because of almost complete conversion of these components.

The reason for this unexpected measured "wrong-way behavior" of the reactor is the delay of the establishment of the new adsorption equilibrium of steam being not considered by the model, which assumes that the steady-state reaction rate at a given temperature is instantaneously reached. Hence, the model only accounts for transient thermal effects, and the desorption kinetics and the adsorption capacity of the catalyst are not considered. So, the overall response time of the reactor of in total 5 min is in about equal amounts the result of the "delayed" ad-/desorption of steam and of the "thermal" behavior. The former effect is not reflected by the model that only considers the steady-state kinetics (only depending on temperature and the gas-phase concentrations of CO, H_2 , and H_2O) and the general transient mass and heat balance.

The transient kinetic experiments presented in Sect. 4.2 clearly show that the reaction rate and the adsorption equilibrium, respectively, require some time to reach steady state, if the steam content in the syngas changes (Fig. 6). The hotspot position at the higher volumetric flow of 60 Lh^{-1} (STP) is located at a reactor length of 5 cm. At this position, the (modeled) CO₂ conversion and, therefore, the content of steam in the syngas is lower (2.4 %) compared to 16 % steam for the lower flow rate (7.5 Lh⁻¹) adjusted before the step change from a low to a high volumetric rate was conducted (subfigure of Fig. 9). Thus, the reaction rate and in turn the level of the hotspot temperature are diminished until the respective surplus of adsorbed steam has desorbed into the gas phase.

Fig. 10 illustrates the result of an experiment conducted in reverse direction, i.e., step change from high to low volumetric flow. Again, the model predicts that the steady state is reached earlier. Now the hotspot temperature slightly moves towards the entrance of the reactor (from l = 5 cm to 4 cm according to the experiment), and a clear but small temperature rise is observed before the new steady state is reached. This is a consequence of the pseudo-homogeneous model used in this work.

At the initially high volume flow, the content of steam at the hotspot position is only 2 %, while at the same reactor position 13 % of steam is present (at steady state). This again leads to the "wrong-way behavior", but now with a higher rate and hotspot temperature, respectively, in the transitional period (0 < t < 120 s). Now a certain additional amount of steam adsorbs in the front region located near the hotspot, which at first has to be produced through the reaction. This takes about 120 s to a higher rate and hotspot temperature in that period compared to the model (not considering this effect).



Figure 10. Hotspot temperature of the wall-cooled fixed-bed reactor after a volume flow step change from 60 to 7.5 L h⁻¹ (STP); inlet composition: 63 % H₂, 15.3 % CO₂, 21.7 % N₂ (methane substitution). Reaction conditions and hotspot positions see Fig. 9. For CO₂ conversion reached at steady state for both volume rates see Fig. 7. The subfigure shows the modeled amount of water at both feed rates at different axial positions in the front part of the reactor at steady state.

5 Conclusions

Transient effects on CO_2 methanation were studied both in an isothermal reactor with regard to kinetic effects as well as in a wall-cooled fixed-bed reactor with regard to additional thermal effects. The steady-state kinetics of CO_2 methanation follow a Langmuir-Hinshelwood approach. The reaction rate is strongly inhibited by steam formed during the reaction.

Isothermal transient kinetic experiments (step change of syngas composition) show a very fast adjustment of the new steady-state reaction rate in case of a step change of the concentrations of CO_2 and H_2 . On the contrary, the steady-state rate is reached with a certain delay (here about 7 min), if the content of steam is changed during a step change from 20 % H_2O to a syngas free of steam. Consequently, the instantaneous establishment of steady-state kinetics is not valid in general.

The overall response time of the wall-cooled methanation reactor (here 5 min) is in about equal amounts the result of the "delayed" ad/desorption of steam and of the "thermal" behavior, if steady-state kinetics (only depending on temperature and the gas-phase concentrations of CO, H_2 , and H_2O) and the general transient mass and heat balance are (only) taken into account.

With regard to practical applications, i.e., for the dynamic operation of a methanation reactor, the overall response time is still short, and the rise in the hotspot temperature is small compared to the steady state. Hence, the danger of a thermal runaway is very low even if the feed rate of the syngas is rapidly changed from a high to a low value.

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Symbols used

с	$[mol m^{-3}]$	molar concentration
c _p	[J kg ⁻¹ K ⁻¹]	thermal capacity
đ	[m]	diameter
$E_{\rm A}$	$[J mol^{-1}]$	activation energy
$\Delta_{ads}H$	$[J \text{ mol}^{-1}]$	adsorption enthalpy
$\Delta_{\rm R} H$	$[kJ mol^{-1}]$	reaction enthalpy
k_0	$[mol(kg s bar)^{-1}]$	reaction rate constant
$K_{0,1}$	[bar ^{-0.5}]	adsorption constant for CO ₂
$K_{0,2}$	[bar ^{-0.5}]	adsorption constant for H ₂
K _{0,3}	$[bar^{-0.25}]$	adsorption constant for H ₂ O
K_p	[-]	equilibrium constant
Nu	[-]	Nusselt number
р	[bar]	partial pressure
r _m	$[mol kg^{-1}s^{-1}]$	reaction rate
R	$[J kg^{-1}mol^{-1}]$	gas constant (8.314)
Т	[K]	absolute temperature
$T_{\rm cool}$	[C]	cooling temperature
$U_{\rm bed}$	$[W m^{-2}K^{-1}]$	overall heat transfer coefficient
u _s	$[m s^{-1}]$	superficial velocity
V	$[m^3s^{-1}]$	volume flow
Χ	[-]	conversion
у	[-]	content (in gas phase)

Greek letters

α	$[W m^{-2}K^{-1}]$	heat transfer coefficient $[m_g^3 m_r^{-3}]$
ε	$[m_{g}^{3}m_{r}^{-3}]$	porosity
λ	$[W m^{-1}K^{-1}]$	thermal conductivity
η	$[\text{kg m}^{-1}\text{s}^{-1}]$	viscosity
ρ	$[\text{kg m}^{-3}]$	density
ν	[-]	stoichiometric factor

Sub- and superscripts

1	CO_2 concerning constants
2	H ₂ concerning constants
3	H ₂ O concerning constants
ax	axial
bed	fixed-bed
eq	equilibrium
g	gas
р	particle
r	reactor
tot	total
w	wall



Abbreviations

LHHW	Langmuir-Hinshelwood-Hougen-Watson
PtG	power-to-gas
CNIC	and the state of the second second

SNG synthetic natural gas

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