Hydrogenation of CO/CO$_2$ mixtures under unsteady-state conditions: Effect of the carbon oxides on the dynamic methanation process

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Abstract

The Power-to-Gas (PtG) process offers the opportunity to store fluctuating renewable energy in form of chemical energy by hydrogenating carbon oxides into methane. In addition, potential carbon point sources often consist of CO/CO$_2$ (CO$_x$) mixtures. Hence, reactor design requires kinetic models valid for unsteady-state operation and a broad spectrum of feed gas compositions. In order to provide the required experimental data basis for derivation of kinetic expressions valid under transient conditions, the dynamic response of a continuously operated fixed-bed methanation reactor is studied by applying periodic step-changes in the feed composition. The obtained results are evaluated based on a simple reactor model, providing the molar flow rate exchanged between the gas bulk and the solid surface for CO, CO$_2$, CH$_4$, and H$_2$O. The results further reveal that the transient kinetic processes at the catalyst surface strongly affect the reactor response under reaction conditions of technical relevance.

Keywords

- Experimental investigation
- Transient behavior
- Step-response
- Simulation based data evaluation
- Methanation
- Periodic experiments
# List of Symbols

| Symbol | Description | Unit |
|--------|-------------|------|
| $d_p$  | Particle size | µm   |
| $F$    | Step response | 1    |
| $f$    | Activity factor | 1    |
| $m_{\text{cat}}$ | Catalyst mass | mg   |
| $N_\Theta$ | Number of limit cycles | 1  |
| $\dot{n}$ | Molar flow rate | mol s\(^{-1}\) |
| $\bar{\dot{n}}$ | Average molar flow rate | mol s\(^{-1}\) |
| $n$    | Molar amount of component | mol |
| $p$    | Operating or partial pressure | bar |
| $R$    | Universal gas constant | J mol\(^{-1}\) K\(^{-1}\) |
| $r$    | Reaction rate | mol kg\(^{-1}\) s\(^{-1}\) |
| $s$    | Cycle split | 1 |
| $T$    | Operating temperature | K |
| $t$    | Time | s |
| $\dot{V}$ | Volumetric flow rate | mL\(_{\text{STP}}\) min\(^{-1}\) |
| $x$    | Molar fraction | 1 |
| $z$    | CO/CO\(_2\) feed ratio | 1 |

*Latin letters*

| Symbol | Description | Unit |
|--------|-------------|------|
| $\alpha$ | Volume variation factor | 1 |
| $\Theta_C$ | Carbon surface coverage | 1 |
| $\nu$    | Stoichiometric coefficient | 1 |
| $\tau$   | Period duration | s |

*Greek letters*

| Symbol | Description | Unit |
|--------|-------------|------|
| $\alpha$ | Volume variation factor | 1 |
| $\Theta_C$ | Carbon surface coverage | 1 |
| $\nu$    | Stoichiometric coefficient | 1 |
| $\tau$   | Period duration | s |

*Subscripts*

| Symbol | Description |
|--------|-------------|
| $+/-$  | Desorbed / re-adsorbed |
| 1, 2   | Initial / final steady-state |
| dil    | Dilution |
| fit    | Fit |
| $i$    | Component $A_i$ |
| in     | Inlet |
| $j$    | Reaction $R_j$ |
| $m$    | Number of limit cycle |
| out    | Outlet |
| RTD    | Residence time distribution |
| ref    | Reference |
| STP    | Standard temperature pressure |
| ss     | Steady-state |
| trans  | Transient |
Abbreviations

| Abbreviation | Description |
|--------------|-------------|
| CO₂          | CO/CO₂      |
| PtG          | Power-to-Gas|
| RTD          | Residence time distribution |

1 Introduction

The Power-to-X process is a promising technology to tackle the challenge of the storage of surplus fluctuating renewable electricity by converting it into chemical energy. During the Power-to-Gas (PtG) process, H₂ is produced from water and renewable electrical energy via electrolysis and subsequently converted with the carbon oxides CO or CO₂ into CH₄ (synthetic natural gas) by the Sabatier reactions (Götz et al., 2016; Rönsch et al., 2016; Kalz et al., 2017). In short and mid-term, the most promising carbon oxide sources are industrial exhaust gases, such as blast-furnace and converter gas from steel industry, gases from cement production or biomass gasification, which potentially consist of CO/CO₂ (COₓ) mixtures of variable composition over time (Sutton et al., 2001; Schöß et al., 2014; Uribe-Soto et al., 2017). Using highly concentrated CO₂ from those sources also offers to reduce industrial CO₂ emissions significantly. In perspective, this technology can be applied to achieve negative emissions, as well, if CO₂ from ambient air is used as reactant (House et al., 2011; Goeppert et al., 2012; Dittmeyer et al., 2019). Such point sources for renewable energy and carbon oxides require local PtG plants in order to avoid costly transport infrastructure and inefficiencies in distribution (Vogt et al., 2019). This also requires plants at a smaller scale tailored to the local availability of the carbon and energy resources. Due to economic reasons such small scale plants, though, probably exhibit small buffer capacities to dampen fluctuations originating from renewable resources (Vogt et al., 2019; Theurich et al., 2020). Consequently, the methanation reactor is supplied by a dynamically changing feed gas mixture and flow rate, which needs the reactor to be operated in an unsteady-state mode. In order to design and operate methanation reactors efficiently and safely under unsteady-state COₓ methanation conditions detailed knowledge on the dynamics of this process upon changes in the H₂/COₓ feed gas composition is necessary.
The exothermic hydrogenation of CO and CO\(_2\) into CH\(_4\) (eqs. (1) and (2)) is performed by means of a porous catalyst, being Ni/Al\(_2\)O\(_3\) the most common compound (Gao et al., 2015). Other metals, such as Co, Ru and Fe are also found to be active and selective, but are only rarely applied commercially so far (Mills and Steffgen, 1974; Gao et al., 2015). The most important side reactions are the reverse water gas shift reaction (RWGS) (eq. (3)), as well as the Boudouard reaction (eq. (4)). The RWGS provides some flexibility towards changes in the feed gas mixture, as it allows to adjust the ratio between CO, CO\(_2\) and H\(_2\) to stoichiometrically favored values. In addition to that the feed gas composition imposes kinetic effects, as well, since CO is known to strongly inhibit the CO\(_2\) methanation process (Inui et al., 1980; Weatherbee and Bartholomew, 1982; Meyer et al., 2020). The Boudouard reaction leads to coke formation, which might cause deactivation of the catalyst. Further minor reactions involved are summarized by Miao et al. (Miao et al., 2016) together with a comprehensive overview of the relevant mechanistic aspects for CO and CO\(_2\) methanation.

\[
\begin{align*}
\text{CO}_2 + 4\text{H}_2 & \rightleftharpoons \text{CH}_4 + 2\text{H}_2\text{O} & \Delta H^{298\text{K}}_R &= -164 \text{ kJ mol}^{-1} \\
\text{CO} + 3\text{H}_2 & \rightleftharpoons \text{CH}_4 + \text{H}_2\text{O} & \Delta H^{298\text{K}}_R &= -205 \text{ kJ mol}^{-1} \\
\text{CO}_2 + \text{H}_2 & \rightleftharpoons \text{CO} + \text{H}_2\text{O} & \Delta H^{298\text{K}}_R &= 41 \text{ kJ mol}^{-1} \\
2 \text{CO} & \rightleftharpoons \text{CO}_2 + \text{C} & \Delta H^{298\text{K}}_R &= -172 \text{ kJ mol}^{-1}
\end{align*}
\]

Dynamically changing feed gas compositions, also designated as concentration forcing, induces an unsteady-state response of the reactor on form of transient behavior of the outlet composition (Bailey et al., 1971; Renken, 1972; Matros, 1987; Marković et al., 2008; Güttel, 2013; Hudgins et al., 2013). Interestingly, different length scales relevant for heterogeneously catalyzed reactions contribute to this response, namely the macro or reactor scale, the meso or pellet scale and the micro scale for processes related to the active surface (Güttel et al., 2020). Most recent studies concentrate on the theoretical investigation of the unsteady-state reactor behavior for CO and CO\(_2\) methanation by means of modeling and simulation. Emphasis is on temperature control (Li et al., 2015; Try et al., 2017; Bremer and Sundmacher, 2019; Kreitz et al., 2019b; Theurich et al., 2020), optimal reactor (Kiewidt and Thöming, 2015; Kreitz et al., 2019b; Fischer and Freund, 2020) or catalyst (Zimmermann et al., 2020) design, and possible performance enhancement (Currie et al., 2018; Nikačević et al., 2020), since the
exothermic methanation is limited by thermodynamic constraints at higher temperatures and puts high demands for safe operation. Those studies, however, rely on reaction kinetics obtained under steady-state conditions, which are not suitable to investigate the reactor behavior under highly dynamic conditions, as will be shown in the present contribution. The reason is that the reactor response is not only governed by residence time distribution, but kinetic processes at the catalyst surface, e.g. ad- and desorption of reacting species, become relevant. In other words, since steady-state kinetics assume equilibrium conditions at the catalyst surface, the respective kinetic processes involved are neglected. In contrast to this assumption, a dynamic and complex system at the catalyst surface arises contributing to the unsteady-state reactor behavior, which requires appropriate consideration of those kinetic processes in the reactor model.

Several experimental studies towards the effect of concentration forcing on the dynamic reactor response for methanation have already been performed at the micro and macro scale (Yadav and Rinker, 1990; Adesina et al., 1995; Klusáček and Stuchlí, 1995; Kreitz et al., 2019a). At the macro scale Stiegler et al. (Stiegler et al., 2019) and Theurich et al. (Theurich et al., 2020) recently demonstrated experimentally that improved temperature control of an unsteady-state methanation reactor is possible in structured reactors system (Stiegler et al., 2019) or by product recirculation (Theurich et al., 2020). With focus at the micro scale one of the first studies was performed by Van Ho and Harriott (Van Ho, 1980) reporting that the CH$_4$ response differs for CO$_2$ and CO methanation. The authors observed a slow exponential decline of CH$_4$ content at the reactor outlet after a switch from a CO$_2$ rich to a CO$_2$ free feed, indicating hydrogenation of an unreactive carbon component during the CO$_2$ free phase. In contrast for CO containing feeds, they reported a rapid increase in the formation rate of CH$_4$, when the CO containing feed is switched off. Underwood and Bennet (Underwood and Bennett, 1984) demonstrated that this behavior is caused by a reactive carbon species adsorbed at the surface. Later, Efstathiou and Bennet (Efstathiou and Bennett, 1989) expanded this work and showed that also an inactive species is hydrogenated to CH$_4$ when the feed is switched from being CO rich to CO free. Moreover, Yadav and Rinker (Yadav and Rinker, 1992) explained that the CH$_4$ behavior after switches from a CO containing to pure H$_2$ feed is mainly determined by the applied H$_2$/CO molar ratio in the CO
rich phase. Fujita et al. (Fujita et al., 1993) conducted diffusive reflectance infrared Fourier-transform spectroscopy (DRIFTS) measurements for the unsteady-state CO and CO₂ methanation and suggested that the highly reactive species only exists in combination with strongly adsorbed bridged CO. Based on these observations different reaction mechanisms were proposed to describe the unsteady-state CO and CO₂ methanation process, which includes the formation of CO as an intermediate species for the CO₂ reaction path (Stuchlý and Klusáček, 1993; Marwood et al., 1994; Aparicio, 1997). In all these studies, however, the CH₄ response is evaluated only. Therefore, no conclusion can be drawn on the interplay of the transient behavior of all reactants detected in the gas phase. In contrast, Bundhoo et al. (Bundhoo et al., 2009) considered all reactants for the unsteady-state CO methanation reaction and provide the dynamic O, H and C mass balances. Based on that they concluded that the availability of metallic sites for hydrogen influences the CH₄ response mostly. However, the unsteady-state methanation reaction was only studied either for CO or CO₂ methanation. The dynamics of the methanation of COₓ mixtures has not been reported so far, neither studied experimentally, nor by modeling and simulation.

Consequently, the present contribution aims at closing the gap between unsteady-state CO and CO₂ methanation experimentally under realistic conditions by applying the novel periodic transient kinetics (PTK) method introduced recently by Meyer et al. (2021). The PTK method is based on the chemical transient kinetics (CTK) method (Raub et al., 2021) and utilizes periodic step-changes between two different feed-gas mixtures in order to induce a periodic transient reactor response. The resulting limit cycle, obtained for each individual component, is statistically analyzed in order to derive an average transient response with improved significance. Furthermore, an internal standard is used to measure the residence time distribution (RTD) in each single experiment in situ. By evaluating the obtained response with a simple unsteady-state reactor model the molar flow rate exchanged between the gas bulk and the solid surface for each individual component is derived as function of time within the limit cycle. As pointed out by Meyer et al. (2021) this methodology provides the basis to separate kinetic processes at the macro scale (e.g., RTD) from those at the micro scale (surface reaction steps) experimentally and is demonstrated to be applicable to realistic reaction conditions. The present
contribution, hence, exploits the capabilities of the PTK method for derivation of mechanistic insights under transient conditions using CO\textsubscript{x} methanation as an example.

In particular, we put emphasis on the influence of the CO/CO\textsubscript{2} ratio at constant H\textsubscript{2}/CO\textsubscript{x} ratio on the transient response. Therefore, a carbon rich feed stream consisting of a particular CO\textsubscript{x} ratio between pure CO and pure CO\textsubscript{2} is exchanged with a carbon free stream by step-shaped periodical switching at the reactor inlet. The transient response of the isothermal reactor is monitored for the most relevant carbonaceous species (i.e., CO, CO\textsubscript{2}, CH\textsubscript{4}) and H\textsubscript{2}O at the outlet. Additionally, an unsteady-state reactor model is applied based on steady-state reaction kinetics developed for the CO\textsubscript{x} methanation reaction (Meyer et al., 2020), in order to evaluate the experimental data. The applied experimental procedure and model-based evaluation enables us to investigate the processes at the catalyst surface without superimposed RTD effects. We are thereby able to link the kinetic processes at the catalyst scale to the unsteady-state reactor behavior. Furthermore, we show that the transient CH\textsubscript{4} response depends not only on the kinetic processes of this species at the catalyst surface but also on the transient behavior of the other reactants. It is further revealed that steady-state reaction kinetics are not sufficient to describe the dynamic reactor behavior for fast changes of the inlet composition.

2 Materials and Methods

2.1 Catalyst

The catalyst based on 5 wt-% Ni/A\textsubscript{2}O\textsubscript{3} with a particle size of 150 – 200 µm is synthesized via an incipient wetness method. By means of H\textsubscript{2} chemisorption and N\textsubscript{2} physisorption measurements (3Flex, Micromeritics) the specific surface area (Brunauer-Emmett-Teller) of 168 m\textsuperscript{2} g\textsuperscript{−1} with an average pore diameter of 8 nm and the H\textsubscript{2} adsorption capacity of 48 µmol g\textsuperscript{−1} are obtained. Assuming spherical Ni nanoparticles as well as an adsorption stoichiometry of two for H\textsubscript{2} a crystallite size of 7.5 nm, a metal surface area of 3.7 m\textsuperscript{2} g\textsuperscript{−1}, and a metal dispersion of 14.2 % is determined. More information about the catalyst synthesis and characterization procedure can be found in our previous publication (Meyer et al., 2020).
2.2 Experimental Setup

For all experiments a stainless-steel fixed bed reactor with an inner diameter of 4.5 mm and a total length of 30 cm is used, in which 50 mg of the catalyst sample diluted with 100 mg of inert material (200 µm, Al₂O₃, Sasol Puralox) is placed in the center of the isothermal zone. For fixation of the packing 1.2 g glass particles (150 – 200 µm) are used above (ca. 5 cm length) and 0.6 g below (ca. 2.5 cm length) the catalyst packing. The whole packing is framed by quartz wool. The reactor is fed by two separate supply lines, which can be automatically switched by means of a 4/2 way-valve (Fitok, BOSS-4C). The continuous supply of the gases is provided from top of the reactor with separate mass flow controllers (EL-FLOW Prestige, Bronkhorst) for each gas and each supply line. The feed gas compositions in both gas lines can be individually adjusted with the reactants H₂ (5.0 purity, MTI), CO/Ar (90 vol.-% CO 3.8 purity in Ar 5.0 purity, Air Liquide) and CO₂ (4.8 purity, MTI), the internal standard Ar (5.0 purity, MTI) and the inert component He (5.0 purity, MTI). Directly at the reactor outlet the gas flow can be diluted with a mixture of H₂ and Ne (both 5.0 purity, MTI) prior entering the analytics. The analytics consist of a gas chromatograph (GC-2010, Shimadzu) and a mass spectrometer (MS, Cirrus 3-XD, MKS). The MS analyzes following species by calibrated mass-to-charge ratios quantitatively: 2 (H₂), 15 (CH₄), 18 (H₂O), 28 (CO and CO₂), 40 (Ar), and 44 (CO₂). The calibration of H₂O is done in situ with pure CO methanation reaction at 556 K, since significant side reactions are proven to be absent under these conditions (Meyer et al., 2020). Additionally, the mass-to-charge ratios for higher hydrocarbons 26 (C₂H₆ and C₃H₈) and 43 (C₃H₈) are measured qualitatively. A flowsheet of the experimental setup is given in the supplementary information (see SI, Figure S1) and a more detailed description of the experimental setup can be found elsewhere (Meyer et al., 2021).

2.3 Experimental Procedure

The experiments are performed at different temperatures and CO₅ feed ratios with the same catalyst sample under the conditions given in Table 1 and Table 2. Prior to the experiments the catalyst is reduced in situ at 673 K for 12 h under flowing H₂ (50 mLSTP min⁻¹). Between each experiment the catalyst was conditioned for 1 h under flowing H₂ (125 mLSTP min⁻¹) at 603 K in order to ensure an
identical initial state, which is verified by reproduction of the first experiment after three consecutive experiments with negligible deviations only (see SI, Figure S2).

Table 1: Operating conditions.

| Parameter                      | Symbol | Unit      | Value |
|--------------------------------|--------|-----------|-------|
| Catalyst mass                 | $m_{\text{cat}}$ | mg        | 50    |
| Particle size                  | $d_p$  | $\mu$m    | 200   |
| Total inlet flow rate          | $\dot{V}_{\text{in,STP}}$ | mL$_{\text{STP}}$ min$^{-1}$ | 250   |
| Dilution flow rate (H$_2$/Ne)  | $\dot{V}_{\text{dil,STP}}$ | mL$_{\text{STP}}$ min$^{-1}$ | 245/5 |
| Operating temperature          | $T$    | K         | 513 – 576 |
| Operating pressure             | $p$    | bar       | 2     |
| Period duration                | $\tau$ | s         | 240   |
| Split                          | $s$    |           | 0.5   |

For concentration forcing experiments the H$_2$/He feed gas mixture is periodically exchanged with the CO$_x$/H$_2$/He/Ar mixture by step-shaped switching between feed line 1 and 2 according to Table 2. In each experiment 50 periods are performed. The respective experiments are carried out for different CO$_x$ ratios as defined by the parameter $z$ (eq. (5)), where $z = 0$ corresponds to pure CO$_2$ methanation and $z = 1$ to pure CO methanation. The half-period after the switch from H$_2$/He to CO$_x$/H$_2$/He/Ar is denominated as build-up phase, while the other half-period is referred to as back-transient phase.

$$z = \frac{p_{\text{CO,in}}}{p_{\text{ref}}} = 1 - \frac{p_{\text{CO}_x,\text{in}}}{p_{\text{ref}}} \quad (5)$$

The reference partial pressure is set constant to $p_{\text{ref}} = 0.2$ bar, which corresponds to the constant CO$_x$ partial pressure in all experiments. Furthermore, the partial pressure of H$_2$ and the CO$_x$/H$_2$ ratio is kept constant in all experiments. Note that the reactor is operated in differential mode according to Shekhtman and Yablonsky (Shekhtman and Yablonsky, 2005) in all experiments by keeping the total carbon oxide conversion below 10%.

Table 2: Inlet partial pressures $p_{i,\text{in}}$ for component $A_i$ in feed line 1 and 2.

| Feed line | $A_i$ | $p_{i,\text{in}}$ / bar | $p_{i,\text{in}}$ / bar |
|-----------|-------|--------------------------|--------------------------|
|           | He    | 0.98                     | 1.20                     |
|           | H$_2$ | 0.80                     | 0.80                     |
Model-based data evaluation

2.4.1 Calculation of molar flow rates

Neon (Ne) is used as external standard by dosing it into the reactor outlet stream with a constant molar flow rate $\dot{n}_{Ne}$. Hence, the measured Ne signal $x_{Ne}$ is only affected by changes in the overall molar flow rate at the outlet $\dot{n}_{out}$ (eq. (6)). By comparison of the measured molar fraction $x_{Ne}$ under reaction conditions with the reference obtained under absence of reaction $x_{Ne,ref}$, parameter $\alpha$ can be derived with eq. (7). The parameter $\alpha$ is used in order to quantify the outlet molar flow rate $\dot{n}_{i,\text{out}}$ from the measured molar fraction $x_{i,\text{out}}$ of component $A_i$ with eq. (8). $\dot{V}_{\text{in,STP}}$ represents the overall inlet volumetric flow rate under standard temperature $T_{\text{STP}}$ and pressure $p_{\text{STP}}$; $R$ is the universal gas constant.

$$x_{Ne}(t) = \frac{\dot{n}_{Ne}}{\dot{n}_{out}(t)}$$  (6)

$$\alpha(t) = \frac{x_{Ne,\text{ref}}}{x_{Ne}(t)} = \frac{\dot{n}_{out}(t)}{\dot{n}_{out,\text{ref}}}$$  (7)

$$\dot{n}_{i,\text{out}}(t) = \alpha(t) x_{i,\text{out}}(t) \frac{\dot{V}_{\text{in,STP}}}{R T_{\text{STP}}}$$  (8)

The evaluation of the transient behavior of the components is performed based on the PTK method, which is described in detail elsewhere (Meyer et al., 2021). Therefore, the outlet molar flow rate $\dot{n}_{i,m}$ of component $A_i$ is measured during consecutive limit cycles $m$ first. Afterwards, the average limit cycle in terms of outlet molar flow rate $\overline{\dot{n}}_{i,\text{out}}$ for all measured compounds is derived by using data for $N_\Theta = 25$ limit cycles according to eq. (9). Figure S3 (see SI) illustrates a representative average limit cycle at 556 K and $z = 0.5$. Due to the very narrow standard deviation interval (shaded areas), we will only show the average values in the following figures for sake of clarity.

$$\overline{\dot{n}}_{i,\text{out}}(t) = \frac{1}{N_\Theta} \sum_{m=1}^{N_\Theta} \dot{n}_{i,m,\text{out}}(t)$$  (9)
The unsteady-state behavior of each component is determined to some extent by transport processes at different length scales. As the internal standard (Ar) does not interact with the catalyst surface, its transient behavior depicts all transport processes on reactor and particle scale. Hence, the internal standard represents the residence time distribution (RTD) of the system. Any deviation from the Ar response, thus, indicates the interaction of the respective component $A_i$ with the catalyst surface. By normalizing the measured outlet molar flow rate $\dot{n}_{i,\text{out}}$ with the values of its initial ($\dot{n}_{i,\text{ss},1}$) and final steady-state ($\dot{n}_{i,\text{ss},2}$), the step response $F_i$ can be derived for each component $A_i$ according to eq. (10).

$$F_i(t) = \frac{\dot{n}_{i,\text{out}}(t) - \dot{n}_{i,\text{out},\text{ss},1}}{\dot{n}_{i,\text{out},\text{ss},2} - \dot{n}_{i,\text{out},\text{ss},1}}$$

Based on the step response of the internal standard $F_{\text{Ar}}$ the expected transient behavior of the outlet molar flow rate of each component $\dot{n}_{i,\text{RTD}}$ governed by RTD of the reactor only can thus be derived according to eq. (11). The deviation of the measured outlet molar flow rate of each component $\dot{n}_{i,\text{out}}$ from the expected RTD during a transient process can be quantified by eq. (12) and is assigned to the transient molar flow rate $\dot{n}_{i,\text{trans}}$. Since the obtained RTD is equal for all species, the transient molar flow rate characterizes processes involved at the catalyst surface. Note, the reference molar flow rates $\dot{n}_{i,\text{out},\text{ss},2}$ are given in the SI (Table S1).

$$\dot{n}_{i,\text{RTD}}(t) = F_{\text{Ar}}(t) (\dot{n}_{i,\text{out},\text{ss},2} - \dot{n}_{i,\text{out},\text{ss},1}) + \dot{n}_{i,\text{out},\text{ss},1}$$

$$\dot{n}_{i,\text{out}}(t) = \dot{n}_{i,\text{RTD}}(t) + \dot{n}_{i,\text{trans}}(t)$$

### 2.4.2 Model description

In order to evaluate the experimental results an unsteady-state, quasi-homogeneous plug-flow reactor model is applied accounting for the change in molar amount with reaction extent (eq. (13)). The molar flow rate $\dot{n}_i$ is assumed to change along the axial direction $x$, due to chemical reaction.

$$\frac{\partial \dot{n}_i}{\partial t} = \frac{\partial \dot{n}_i}{\partial x} \text{d}x + \text{d}m_{\text{cat}} (v_{i,\text{CO}} r_{\text{CO}} + v_{i,\text{CO}_2} r_{\text{CO}_2})$$

The reaction kinetics used to calculate the mass specific reaction rate $r_j$ for CO and CO$_2$ methanation are taken from our previous study, which is performed with the same catalyst (Meyer et al., 2020) and given in the SI (eq. (S1) and (S2)). It is important to emphasize that the model is thus not capable to
predict dynamic processes at the catalyst surface. Therefore, any transient process on the catalyst scale causes a deviation between simulated and measured data, which will be evaluated in this study. The kinetic parameters are taken from Meyer et al. (Meyer et al., 2020) and scaled with activity factors $f_j$ according to the experimental results obtained at 556 K (see SI, Table S2). The equation further requires the stoichiometric coefficients $v_{i,j}$ and the catalyst mass $m_{cat}$. Furthermore, isothermal and isobaric conditions as well the absence of internal and external transport limitations are assumed, which was verified previously (Meyer et al., 2020). The model equations are axially discretized (50 nodes) with the method of lines and solved with the ode45 solver (explicit Runge-Kutta method) using default parameters in Matlab (Version R2017b).

Importantly, the signal at the reactor inlet is not ideal, due to the flow pattern in the piping between the 4/2-way valve and the reactor inlet. Therefore, we measured the step response at the reactor inlet ($x = 0$) by MS using Ar as tracer and applied polynomial fitting to derive the cumulative function of the RTD at the reactor inlet $F_{fit}$ (results see SI, Figure S4). The measured step response was divided in three phases for simplification and according to the expected real laminar flow behavior (eq. (14)). It comprises an induction phase prior to $t_1$ and a steady-state beyond $t_2$. The transient phase between those boundaries is fitted. The values for $t_1 = 0.3$ s and $t_2 = 8$ s are defined ($F_{fit}(t_1) = 0.01$, $F_{fit}(t_2) = 0.99$) in order to neglect the influence of noise of the measured signal. This approach was used in order to calculate the molar flow rate at the reactor inlet $\dot{n}_i$ from the ideal step assumed for $\dot{n}_{i,in}$ (eq. (15)).

$$F_{Ar,0}(t) = \begin{cases} 0 & t < t_1 \\ F_{fit}(t) & t_1 \leq t \leq t_2 \\ 1 & t > t_2 \end{cases} \quad (14)$$

$$\dot{n}_i(x = 0, t) = F_{Ar,0}(t) \dot{n}_{i,in} \quad (15)$$

3 Results and Discussion

This section is structured as follows. The results obtained for relevant reactants (CO and CO$_2$) and products (H$_2$O and CH$_4$) are presented in that order. For each component the build-up and back-
transient phase is reported consecutively. In order to avoid repetition the results are discussed in context of that sequence, as far as possible.

3.1 Build-up behavior of CO

Figure 1 (left) shows the step response for CO within the first 40 s of the build-up phase for CO feed containing compositions at different temperatures. Note, that the trend in CO for pure CO\textsubscript{2} methanation ($z = 0$) is not shown, as the amount of CO is close to the lower detection limit under the applied conditions, which is in accordance with literature (Gao et al., 2012; Koschany et al., 2016). After switching from a CO free to a CO rich feed an induction phase is observed within the first 5 s, which is caused by the RTD measured \textit{in situ} and indicated by the dash-dotted line. In the following the CO step response increases approaching steady-state levels asymptotically. While the CO step response follows the RTD with small deviation for $z \geq 0.5$, the deviations observed for $z = 0.1$ will be discussed in the following.

Figure 1 (right) shows the transient molar flow rates, representing the difference between the expectations from RTD and the measured absolute molar flow rates. Negative values indicate a CO sink with respect to the gas phase, while positive values correspond to a source. Since adsorption of CO at the active surface and the conversion by chemical reaction are the only kinetic processes not covered by the RTD of the internal standard, CO is likely to be stored at the active surface by adsorption and converted by subsequent surface reactions. To make it more precise: We assume that the internal standard is chemically inert and capable of describing all kinetic processes related to pure transport of CO (e.g. pore diffusion), adequately. Therefore, any difference in RTD between the internal standard and CO can directly be associated to surface related processes, being the CO adsorption at the active surface and subsequent conversion the most important during the build-up phase. The simulated transient molar flow rates, depicted in grey, are obtained assuming steady-state reaction kinetics as introduced above. Therefore, adsorption processes of CO at the active surface are not covered by the simulation results.
Figure 1: Left: Step response of CO (colored lines) and Ar (RTD, dash-dotted line); right: transient molar flow rates of CO for different CO feed ratios $z$ and temperatures during the build-up phase; the simulated transient molar flow rates are shown in grey.

The measured transient molar flow rate first decreases more strongly than predicted by the simulation, which is equivalent with an increase in the stored amount of CO at the catalyst surface. Note, that the results of CO, only, allow no conclusion whether CO is adsorbed or if it is even converted. Following a minimum, the transient flow rate is approaching zero again and thus steady-state. The minimum is more pronounced than simulated and shows that the driving force for CO adsorption at the catalyst surface is high in the early stage of the build-up phase, where the surface coverage by CO is still low. With increasing CO surface coverage, the adsorption rate decreases and thus the transient molar flow rate approaches zero, indicating that the CO surface coverage is in steady-state. Furthermore,
significant deviations with respect to the temperature dependency are observed. Obviously, the simulated transient molar flow rates of CO predict a sink being more pronounced at higher temperatures. This trend is reasonable, since the hydrogenation rate of CO increases with temperature. In contrast, adsorption is favored at lower temperatures, which compensates decreasing reaction rates to a certain extent, leading to the small effective impact of temperature on the measured transient molar flow rates.

Figure 1 (right) also shows that the absolute value of the minimum transient molar flow rate is rather constant in the investigated temperature range, while a clear trend can be observed related to the $z$ values. In particular, the minimum is expressed more significantly for higher CO fractions in the feed gas and the integrated transient molar flow rate over time increases with the CO fraction, as well. The significant effect of the CO concentration on CO adsorption and the absence of a temperature effect agrees with literature (Polizzotti and Schwarz, 1982; Biloen et al., 1983; Alstrup, 1995). For example, Polizotti et al. (Polizzotti and Schwarz, 1982) stated that the catalyst surface remains nearly saturated with CO irrespective of the temperature in the range between 500 and 700 K. In contrast the amount of adsorbed CO depends clearly on its partial pressure (Bartholomew and Pannell, 1980), which leads to a decrease of the surface coverage of CO with decreasing CO partial pressure. The results also correspond to our findings for simultaneous CO$_x$ methanation under steady-state conditions exhibiting exclusive CO hydrogenation for $z > 0.1$, even though CO$_2$ is present (Meyer et al., 2020). Under those conditions, CO occupies essentially all sorption sites, which effectively prevents CO$_2$ from adsorption (Meyer et al., 2020).

3.2 Back-transient behavior of CO

Figure 2 shows the results obtained during the back-transient phase of CO after switching from CO rich to CO free feed gas. The step response is provided in the SI (Figure S5) and show no significant difference to the RTD irrespective of feed gas compositions and temperatures. Based on the transient molar flow rates, however, a dependence on temperature and CO$_x$ ratio can be observed. At first the transient molar flow rate is positive, indicating CO desorption from the surface into the gas phase. In
the following, the transient molar flow rate drops for all CO containing feeds to negative values before approaching zero in the steady-state. The negative transient molar flow rates suggest that CO from the gas phase adsorbs again at sorption sites just released by conversion into CH₄. Therefore, the desorption step appears to be consecutively followed by re-adsorption and subsequent conversion during the back-transient phase.

![Figure 2: Transient molar flow rates of CO for different COₓ feed ratios z and temperatures during the back-transient phase; the simulated transient molar flow rates are shown in grey.](image)

Furthermore, a significant effect of the COₓ feed ratio and the temperature is apparent from the measured as well as the simulated transient molar flow rates both agreeing with each other with respect to the trends. For a decreasing CO content in the feed gas mixture the transient molar flow rate diminishes, since the reservoir of adsorbed CO becomes smaller, and the reported effects are thus less expressed. The decrease in temperature, interestingly, affects the intensity of the desorption and re-adsorption step. This becomes apparent by comparing the integrated values of the transient molar flow rate distinguished between the desorption (positive values) and re-adsorption (negative values) phase (Table 3). For $z \geq 0.5$ the desorbed molar amount of CO decreases with increasing temperature,
while the re-adsorbed amount increases. This can be explained by the increasing reaction rate with temperature, which leads to fast conversion of adsorbed CO species and therefore more available sorption sites during the transient phase. Consequently, desorption is less likely, while re-adsorption is favored. For \( z = 0 \) no clear trend can be observed, since quantification is difficult regarding the precision of measurement. These findings are supported by literature, as Underwood et al. have shown for pure CO methanation that a small amount of CO desorbs, indeed (Underwood and Bennett, 1984). Furthermore, this might also indicate that CO adsorbs at the catalyst surface without being transformed into a reactive carbon species at lower temperatures, which is in accordance to Agnelli et al. (Agnelli et al., 1998). The re-adsorption during the back-transient phase, however, is not yet reported in scientific literature to the best of our knowledge.

Table 3: Total molar amount of CO desorbed \((n_{\text{trans}},+)\) and re-adsorbed \((n_{\text{trans}},-)\) during the back-transient phase obtained by integration over the respective time interval.

| molar amount of CO / \(\mu\text{mol}\) | 576 K | 556 K | 533 K | 513 K |
|---------------------------------------|-------|-------|-------|-------|
| \( z / 1 \)  | \( n_{\text{trans}},+ \) | \( n_{\text{trans}},- \) | \( n_{\text{trans}},+ \) | \( n_{\text{trans}},- \) | \( n_{\text{trans}},+ \) | \( n_{\text{trans}},- \) |
| 1.0 | 0.5 | 7.2 | 0.8 | 5.8 | 1.7 | 4.1 | 2.2 | 2.2 |
| 0.5 | 0.1 | 3.7 | 0.2 | 3.3 | 0.4 | 2.9 | 0.8 | 2.0 |
| 0.1 | 0.0 | 0.0 | 0.0 | 0.5 | 0.0 | 0.8 | 0.2 | 0.3 |

3.3 Build-up behavior of \( \text{CO}_2 \)

Figure 3 shows the transient molar flow rates for \( \text{CO}_2 \) with the corresponding step response provided in the SI (Figure S6). The deviation of the observed step response of \( \text{CO}_2 \) from the RTD for pure CO methanation conditions \((z = 1)\) indicates that the formation of \( \text{CO}_2 \) is the result of a slow secondary reaction path. This is supported by Zarfl et al. (Zarfl et al., 2015), where the Boudouard and/or the WGS reaction are denoted as the most likely reaction pathways (Kopyscinski et al., 2010; Gao et al., 2012). The transient molar flow rates are close to zero, though, since the \( \text{CO}_2 \) formation under steady-state conditions is negligible (see SI, Table S1).
The negative transient molar flow rates indicate interaction of CO\textsubscript{2} with the solid surface similar to CO, exhibiting the minimum early during the build-up phase, as well. The amount of adsorbed CO\textsubscript{2}, expressed by the minimum and the integrated transient molar flow rate, is reduced by increasing the fraction of CO in the feed (raising $z$). For an equimolar mixture of CO and CO\textsubscript{2}, for instance, only a small minimum is observed, which suggests a mainly inert behavior of CO\textsubscript{2} at high CO concentrations. This agrees with several studies performed under steady-state conditions (Van Herwijnen, 1973; Polizzotti and Schwarz, 1982; Weatherbee and Bartholomew, 1982; Biloen et al., 1983; Alstrup, 1995; Falbo et al., 2019; Meyer et al., 2020). In particular, CO adsorption is favored compared to the other species, which is even more pronounced at high CO concentrations. Therefore, CO occupies nearly all sorption sites, preventing CO\textsubscript{2} from adsorption, which consequently acts rather inert also with respect to the hydrogenation reaction. The negative transient molar flow rate at $z = 0.5$, however, might be caused by non-competing adsorption of CO\textsubscript{2} on the support being Al\textsubscript{2}O\textsubscript{3} in the present study (Friedland et al., 2020). While Al\textsubscript{2}O\textsubscript{3} is known for its potential to chemisorb CO\textsubscript{2} (Manchado et al., 1994), the adsorption
potential of CO is low (Manchado et al., 1994) and thus assumed negligible. The deviation between measured and simulated transient molar flow rates confirms that CO indeed adsorbs on the surface.

3.4 Back-transient behavior of CO$_2$

The step response (Figure 4, left) shows no significant deviation of the CO$_2$ back-transient behavior from the RTD except for pure CO hydrogenation ($z = 1$). Based on the evaluation of the transient molar flow rates, however, an interaction with the catalyst surface is evident for $z < 1$ (Figure 4, right). Similar to CO, a positive transient molar flow rate of CO$_2$ is observed during the early stage, indicating desorption of previously adsorbed CO$_2$ into the gas phase. The amount of desorbed CO$_2$ is decreasing strongly with raising temperature (Table 4), which is probably weakly bound on the surface (Falconer and Zağli, 1980; Friedland et al., 2020). In the following, the transient molar flow rate drops as more CO$_2$ is converted, which agrees with the simulated values at higher temperatures regarding the principle trend. The quantitative deviation between the measured and simulated transient molar flow rates is caused by the sorption processes, not accounted for in the steady-state reaction kinetics. At high temperatures the transient molar flow rate even reaches negative values, which indicates re-adsorption of CO$_2$ in the back-transient phase similar to the respective discussion for CO. In contrast to the steady-state results conversion of CO$_2$ can be detected even for high amounts of CO in the feed gas mixture (e.g. $z = 0.5$) (Meyer et al., 2020). This can be explained by the simultaneous decrease of CO and CO$_2$ in the gas phase during the back-transient half-period. Therefore, the dominating CO adsorption at the surface becomes less pronounced over time, which provides more sorption sites available for CO$_2$ to adsorb and subsequently to be converted to CH$_4$.

In the case of pure CO methanation ($z = 1$) the CO$_2$ step response exhibits a peak, whose temporal occurrence and maximum is temperature sensitive. Note, that the trends are displayed in the normalized form and that the corresponding absolute amount of produced CO$_2$ is small. Nevertheless, this peak most likely results from a reaction process or desorption from the support, as the CO$_2$ gas phase concentration is low and thus CO$_2$ should desorb instantaneously after formation under given conditions (Falconer and Zağli, 1980; Friedland et al., 2020). We assume that a particular reaction
mechanism is responsible for this observation, where oxygen containing species stored at the catalyst surface are converted. It has to be stated, however, that further experiments are required to fully understand the underlying reaction mechanism to this observation. Even though, such behavior for CO$_2$ under unsteady-state CO methanation conditions is not yet reported, Mutz et al. (Mutz et al., 2017) suggest different possible oxygen sources. In particular, CO$_2$ could either be generated by a reaction intermediate containing oxygen, the WGS reaction, or directly from the catalyst (e.g. from the lattice).

Figure 4: Left: Step response of CO$_2$ (colored lines) and Ar (RTD, dash-dotted line); right: transient molar flow rates of CO$_2$ for different CO$_x$ feed ratios $z$ and temperatures during the back-transient phase; the simulated transient molar flow rates are shown in grey.
Table 4: Total molar amount of CO\textsubscript{2} desorbed ($n_{\text{trans,+}}$) and re-adsorbed ($n_{\text{trans,-}}$) during back-transient phase obtained by integration over the respective time interval.

| z / 1 | $n_{\text{trans,+}}$ | $n_{\text{trans,-}}$ | $n_{\text{trans,+}}$ | $n_{\text{trans,-}}$ | $n_{\text{trans,+}}$ | $n_{\text{trans,-}}$ |
|-------|----------------------|----------------------|----------------------|----------------------|----------------------|----------------------|
| 1.0   | 0.3                  | 0.0                  | 0.3                  | 0.0                  | 0.1                  | 0.0                  |
| 0.5   | 0.4                  | 2.3                  | 0.5                  | 1.0                  | 0.8                  | 0.2                  |
| 0.1   | 0.1                  | 3.9                  | 0.2                  | 1.7                  | 0.7                  | 0.4                  |
| 0.0   | 0.0                  | 2.4                  | 0.7                  | 0.7                  | 2.5                  | 0.1                  |

3.5 Build-up behavior of H\textsubscript{2}O

In contrast to the previously discussed reactants CO and CO\textsubscript{2} the product H\textsubscript{2}O is formed stoichiometrically by chemical reaction as an adsorbed species, which subsequently desorbs into the gas phase (Miao et al., 2016). The transient molar flow rates of H\textsubscript{2}O during the build-up phase show negative values during the complete half-period (Figure 5) corresponding to a temporal delay from the RTD (see step response in SI, Figure S7), which is most pronounced for low temperature and high CO content in the feed gas. Moreover, the measured transient molar flow rates are negative for all investigated conditions, whereas the simulated results exhibit positive values. This fundamental deviation can be explained by the storage of H\textsubscript{2}O at the catalyst surface, especially by adsorption at Al\textsubscript{2}O\textsubscript{3} sites (Morimoto et al., 1971), which buffers the H\textsubscript{2}O response and is not considered in the model. Furthermore, the surface adsorption can also explain that the strongest temporal delay in the transient molar flow rates is observed for pure CO methanation ($z = 1$), since the H\textsubscript{2}O production rate is decreasing with increasing CO fraction in the feed, due to stoichiometric reasons. Hence, more time is required to fill the H\textsubscript{2}O storage capacity for smaller H\textsubscript{2}O formation rates and therefore for increasing values of $z$. Since the sorption capacity decreases and the H\textsubscript{2}O formation rate increases with raising temperature, H\textsubscript{2}O appears earlier in the gas phase and therefore the deviation from RTD becomes smaller. Our results, however, seem to contradict earlier studies by Klusáček and Stuchlý (Klusáček and Stuchlý, 1995) and Fujita et al. (Fujita et al., 1987), where a rapid overshoot in the H\textsubscript{2}O response is
reported. We attribute those observations to the lower storage capacity of H$_2$O for the catalysts used: Klusáček and Stuchlý (Klusáček and Stuchlý, 1995) used Ni/SiO$_2$, while Fujita et al. (Fujita et al., 1987) pure Ni for their experiments. In particular, it is well-known that the sorption capacity of H$_2$O at SiO$_2$ surfaces is significantly smaller than at Al$_2$O$_3$ (Morimoto et al., 1971), which therefore renders the observed discrepancies between our results and literature to be plausible.

![Graph showing transient molar flow rates of H$_2$O](image)

**Figure 5:** Transient molar flow rates of H$_2$O for different CO$_x$ feed ratios $z$ and temperatures during the build-up phase; the simulated transient molar flow rates are shown in grey.

### 3.6 Back-transient behavior of H$_2$O

Figure 6 shows the back-transient behavior of H$_2$O, which exhibits positive transient molar flow rates throughout the half-period for both the experimental and simulation results (for step response see SI, Figure S8). This can be explained by the decreasing CO$_x$ fraction in the gas phase during the back-transient phase and thus the increasing H$_2$/CO$_x$ ratio with time. Therefore, the H$_2$/CO$_x$ ratio passes kinetically favored values for the CH$_4$ and H$_2$O formation, which causes the positive transient molar flow rates (Meyer et al., 2020). The simulated profiles exhibit a narrow distribution in the early stage of the back-transient phase, while the measured values also show significant amounts of H$_2$O.
appearing in the gas phase in a later stage. The steady-state kinetics used for the simulations account for competitive adsorption of the involved species and the implications on the reaction rate, assuming steady-state, though. Hence, the simulated transient molar flow rates exhibit an increasing H$_2$O release with increasing $z$ values, since the positive effect of high H$_2$/CO$_x$ ratios is most pronounced for those cases (Meyer et al., 2020). The tailing in the H$_2$O signal in the later stage, therefore, is most likely caused by slow desorption of H$_2$O stored at the alumina surface, which is not included in the model. Alternatively, hydrogenation of intermediate formate species (COOH) present at the catalyst surface, which are postulated in literature to be present with a significant amount at the surface under steady- and unsteady-state CO and CO$_2$ methanation conditions (Fujita et al., 1991; Marwood et al., 1994; Falbo et al., 2019; Burger et al., 2020), could cause formation and desorption of H$_2$O and thereby effect the later stage of the back-transient behavior. The profiles for all CO$_x$ feed gas mixtures become similar asymptotically, indicating that it is determined by the same physical processes.

Figure 6: Transient molar flow rates of H$_2$O for different CO$_x$ feed ratios $z$ and temperatures during the back-transient phase; the simulated transient molar flow rates are shown in grey.
While the measured profiles are almost identical for all CO\textsubscript{x} ratios at 576 K, a deviation develops at lower temperatures. For the case of pure CO\textsubscript{2} (z = 0) the transient molar flow rate exhibits a maximum in the early stage followed by an exponential decay, indicating that the desorption of H\textsubscript{2}O is the only relevant process. For CO containing feed gas mixtures, though, a maximum is visible in a later stage of the back-transient phase, which is prominent for 513 K and pure CO methanation (z = 1). This maximum shifts backwards in time with increasing temperature and finally coincides with the maximum for pure CO\textsubscript{2} methanation at 576 K. The temperature induced temporal shift of the maximum for CO methanation (z = 1) suggests that a thermally activated reactive process leads to the occurrence of H\textsubscript{2}O in the gas phase. The observation of two different maxima also indicates that two different processes are responsible for the H\textsubscript{2}O back-transient behavior in case of CO containing feed gas mixtures, whereas for pure CO\textsubscript{2} methanation only one process seems to be responsible. In particular, we assume that H\textsubscript{2}O desorption from the solid surface is occurring irrespective from the CO\textsubscript{x} ratio, while intermediate species adsorbed at the surface are only formed at higher CO partial pressures. This is plausible, since the formation of H\textsubscript{2}O and its adsorption depends on methanation stoichiometry and surface chemistry of the solids only, but is independent on the CO content in the gas phase. The presence and type of intermediate surface species, however, strongly depend on the reaction mechanism governed by the partial pressures of CO, CO\textsubscript{2} and H\textsubscript{2} (Fujita et al., 1993; Miao et al., 2016).

3.7 Build-up behavior of CH\textsubscript{4}

The step response of CH\textsubscript{4} in the build-up phase exhibits an overshoot in the CH\textsubscript{4} signal and higher values than predicted from pure RTD (Figure 7). These observations are in particular depending on temperature, while the trend is similar for all z values. The corresponding measured transient molar flow rates are negative initially but turn to positive values during the build-up phase. The negative values indicate a delayed release of CH\textsubscript{4} compared to the RTD, due to the kinetics of the consecutive formation and desorption steps, while positive values indicate an accelerated CH\textsubscript{4} release into the gas phase. The simulated transient molar flow rates are always positive and immediately appear according
to the RTD. Similar to the discussion for the H₂O signal in the back-transient phase the positive values can be explained by the passing of kinetically favored H₂/CO ratios during the build-up. The measured profiles also exhibit positive transient molar flow rates, since the catalyst surface is initially saturated by H₂, due to the previous back-transient phase under H₂/He flow. This favors the CH₄ formation kinetically, as well. In addition, the dynamics of competitive adsorption of the involved species and their impact on the reaction kinetics have to be considered. In particular, CO is capable of displacing H₂ from the surface, which leads to a reduction of the methanation rate for high CO partial pressures (Underwood and Bennett, 1984; Bundhoo et al., 2009). Other studies emphasize the inhibiting effect of CO on the CH₄ methanation, as well (Goodman et al., 1980; Stuchlý and Klusáček, 1993; Aparicio, 1997). Since, the observed overshoot is smaller and delayed compared to the simulated one, it is most probably caused by passing through an optimal ratio of the adsorbed species, during the transient change of the gas phase composition.

It has to be mentioned that the sequence of the measured maximum transient molar flow rate changes with respect to COₓ ratio over the temperature range. At high temperatures the observed maximum is higher with increasing CO content in the feed gas mixture, while the opposite holds for lower temperatures. According to literature a change in the kinetic regime at temperatures of ca. 490 to 510 K might be responsible for this observation (Goodman et al., 1980; Polizzotti and Schwarz, 1982; Friedland, 2018). Goodman et al. (Goodman et al., 1980) show in their study that at temperatures around 500 K the reaction rate is nearly independent on the pressure, whereas at higher temperatures a clear pressure dependency can be seen. They concluded that at higher temperatures the amount of carbon at the catalyst surface can be decreased by an increasing H₂ partial pressure, which is not possible at lower temperatures where CO is strongly adsorbed (Agnelli et al., 1998). Furthermore, increasing the CO₂ content in the feed gas composition reduces the overshoot in the CH₄ production during the build-up phase due to the decreasing inhibition by CO, which is also predicted by the steady-state model. Aparicio observed the same effect for the variation of the CO partial pressure for the pure CO methanation (Aparicio, 1997), stating that with a decrease of CO in the gas phase the CO coverage
cannot longer approach unity, which dampens the inhibiting effect. In contrast to our results, they observed the opposite temperature induced effect, as the overshoot in CH\textsubscript{4} weakens with increasing temperature. However, they conducted their study with a large excess of H\textsubscript{2}, which might explain the difference, as our results are in accordance with Bundhoo et al. (Bundhoo et al., 2009) and Stuchlý and Klusáček (Stuchlý and Klusáček, 1993).

![Figure 7](image)

Figure 7: Left: Step response of CH\textsubscript{4} (colored lines) and Ar (RTD, dash-dotted line); right: transient molar flow rates of CH\textsubscript{4} for different CO\textsubscript{x} feed ratios \( z \) and temperatures during the build-up phase; the simulated transient molar flow rates are shown in grey.

Interestingly, the temporal occurrence of the maximum in the transient molar flow rate of CH\textsubscript{4} nearly coincides with the minimum in H\textsubscript{2}O, as Figure 8 shows exemplarily for different temperatures and \( z = \)
Since both CH₄ and H₂O are formed stoichiometrically, this observation suggests that H₂O is adsorbed at the surface, while CH₄ desorbs almost immediately after being formed. Hence, the presence of adsorbed H₂O species does not hamper the CH₄ formation drastically, while H₂O in the gas phase is reported to inhibit the CH₄ formation rate (Marwood et al., 1994; Borgschulte et al., 2013). It has to be mentioned that the observation strongly depends on the catalyst support used, since Al₂O₃ provides a significant sorption capacity for H₂O in contrast to SiO₂. Klusáček and Stuchlý (Klusáček and Stuchlý, 1995) as well as Fujita et al. (Fujita et al., 1987), for instance, report a slow increase in the CH₄ and a fast increase in the H₂O signal. Both studies used either SiO₂ as support material (Klusáček and Stuchlý, 1995) or pure Ni (Fujita et al., 1987), both known to provide a lower storage capacity for H₂O compared to Al₂O₃ used in the present work.

Figure 8: CH₄ and H₂O transient molar flow rates for z = 0.1 at different temperatures during build-up phase.

3.8 Back-transient behavior of CH₄

Figure 9 (left) shows that a switch from a COₓ rich to a COₓ free feed gas is accompanied with a positive deviation of the CH₄ step response compared to the RTD with a distinct maximum for CO containing feed gas mixtures. The maximum increases with decreasing temperature by a factor of up to 3.5 with respect to the steady-state value. This indicates the high sensitivity of the system towards an exchange of COₓ rich with a COₓ free feed especially for low temperatures and high z values and agrees with
literature (Bundhoo et al., 2009). Furthermore, the maximum decreases with decreasing $z$ and is not observed for $z = 0$ as expected for pure CO$_2$ methanation conditions (Van Ho, 1980; Fujita et al., 1991; Marwood et al., 1994).

The corresponding transient molar flow rate is positive throughout the complete back-transient phase (Figure 9, right). The duration of the process as well as the total amount of released CH$_4$ thereby depends strongly on temperature and CO$_x$ feed ratio (Table 5). At higher temperatures, the process is completed faster and more CH$_4$ is produced, while higher $z$ values also lead to higher amounts of released CH$_4$. In all cases the amount of released CH$_4$ predicted by simulation is significantly lower than the measured values, while the predicted duration of CH$_4$ appearance is shorter and independent on temperature. This indicates that additional carbonaceous species are stored at the catalyst surface during the dynamic experiments, which are converted into CH$_4$ under transient conditions and thereby increase the CH$_4$ signal above the values predicted from steady-state kinetics. In contrast to pure CO$_2$ feed gas mixtures for CO containing feeds ($z > 0$) the form of the CH$_4$ transient molar flow rate changes with temperature. At 576 K the CH$_4$ profile exhibits one maximum and an exponential decay, whereas with decreasing temperature the peak broadens and a shoulder appears for high fractions of CO ($z \geq 0.5$). This indicates that two species at the catalyst surface are hydrogenated to CH$_4$ under CO$_x$ methanation conditions, whereas only one seems to be converted in case of pure CO$_2$ methanation. The first species is immediately converted leading to the appearance of the early maximum, which occurs at ca. 7 s for all cases. The temporal occurrence of this first maximum is predicted correctly by the simulations assuming steady-state kinetics. This underlines that the highly reactive intermediate is hydrogenated immediately without preceding dynamic effects at the catalyst surface, which are neglected by the model. The hydrogenation of the second species is temperature dependent and appears as the shoulder following the maximum in particular at 513 K. Similar to the discussion of the back-transient phase for H$_2$O the second maximum, pronounced as a shoulder here, shifts to earlier times with increasing temperature and finally coincides with the first maximum.
Figure 9: Left: Step response of CH4 (colored lines) and Ar (RTD, dash-dotted line); right: transient molar flow rates of CH4 for different COx feed ratios z and temperatures during the back-transient phase; the simulated transient molar flow rates are shown in grey.

The presence of highly reactive intermediates adsorbed at the surface is supported by literature reports, which suggest that a solely carbon containing species, denoted as Cα, under steady-state (Galuszka et al; Alstrup, 1995; Sehested et al., 2005) and unsteady-state (Underwood and Bennett, 1984; Efstatthiou and Bennett, 1989; Fujita et al., 1991; Fujita et al., 1993; Fujita and Takezawa, 1997) methanation conditions exists. Fujita et al. (Fujita et al., 1993) conducted DRIFTS measurements for transient CO and CO2 hydrogenation experiments and show that the highly reactive species is not present under CO2 methanation conditions, which is consistent with our results. The authors observed
mainly bridged adsorbed CO along with a formate species at the catalyst surface during the CO₂ hydrogenation, whereas a considerable amount of linear adsorbed CO and Cₓ exists for CO methanation. It is concluded that the linear adsorbed CO retards the hydrogenation of Cₓ, but disappears rapidly after switching to pure H₂, which allows a fast hydrogenation of Cₓ subsequently. According to Yadav and Rinker (Yadav and Rinker, 1992) the amount of atomic carbon stored at the surface increases with increasing CO/H₂ molar ratio, which is consistent with our results, since the maximum in CH₄ signal is reduced for low z values and thus low CO/H₂ ratios. The chemical nature of the reactive species, however, being pure carbon or not, is still under debate. Recent studies proposed kinetic rate equations for CO₂ and even COₓ methanation via hydrogen assisted C-O bond cleavage steps towards COH intermediates (Underwood and Bennett, 1984; Koschany et al., 2016; Falbo et al., 2019; Burger et al., 2020), whereas Burger et al. (Burger et al., 2020) found no evidence of elemental carbon species even under CO methanation conditions. Importantly, the re-adsorption of CO and CO₂ observed in the back-transient phase most probably also contribute to the formation of excess CH₄ in addition to the carbonaceous species adsorbed at the catalyst surface. This was often neglected in the past (Van Ho, 1980; Underwood and Bennett, 1984; Yadav and Rinker, 1992; Fujita et al., 1993; Stuchlí and Klusáček, 1993), probably because re-adsorption was not observed so far, due to experimental limitations.

Table 5: Total molar amount of CH₄ desorbed (n_{trans,+}) during back-transient phase obtained by integration over the respective time interval.

| z / 1 | 576 K / n_{trans,+} | 556 K / n_{trans,+} | 533 K / n_{trans,+} | 513 K / n_{trans,+} |
|-------|---------------------|---------------------|---------------------|---------------------|
| 1.0   | 15.0                | 13.8                | 11.0                | 9.0                 |
| 0.5   | 12.9                | 11.4                | 9.0                 | 7.6                 |
| 0.1   | 10.3                | 8.7                 | 6.6                 | 5.6                 |
| 0.0   | 7.5                 | 4.4                 | 3.9                 | 2.5                 |
4 Conclusion

The influence of CO, CO₂ and mixtures thereof on the transient behavior of the reactants during methanation was investigated experimentally for a Ni/Al₂O₃ catalyst in a continuously operated fixed-bed reactor under technical relevant conditions by means of the novel PTK method. Therefore, periodic step changes are performed at the reactor inlet between CO₂ rich and CO₂ free gas compositions and vice versa, in order to study the build-up and back-transient half-period. For all reactants the temporal profiles of the periodic step response are obtained and deconvoluted from the RTD measured in situ. Moreover, an unsteady-state reactor model is developed based on steady-state COₓ methanation kinetics. This sophisticated approach combining experiments and simulations was used to deduce the transient behavior of the kinetic processes taking place at the solid catalyst surface and their impact on the observed unsteady-state reactor behavior. From the obtained results the following conclusions can be drawn:

1. A reactor model based on unsteady-state material balances and steady-state reaction kinetics is not capable to predict the transient behavior of the reactor, neither during the build-up nor in the back-transient phase. The kinetic processes taking place at the catalyst surface are strongly affecting the reactor response, instead, and need to be considered at least for step changes at the reactor inlet. In addition, the storage capacities of the species by adsorption at the solid surface play an important role for the reactor response.

2. The experimental results obtained for CO, CO₂, CH₄ and H₂O during build-up and back-transient phase are consistent and agree with literature reports in principle. In particular, the support provides significant storage capacity for adsorbed H₂O and CO is found to strongly adsorb at the active material and thereby inhibits CO₂ hydrogenation. In addition to the state-of-knowledge, CO and CO₂ are found to re-adsorb at the catalyst surface during the back-transient phase and are partially converted into CH₄ and H₂O.

3. The dynamic reactor behavior is strongly influenced by the kinetic processes occurring at the solid surface, which are governed by the COₓ and the H₂/COₓ ratio, as well as the reaction temperature.
In particular, maxima in the transient signal of CH$_4$ compared to the steady-state values are observed during both the build-up and back-transient phase, induced by passing kinetically favored gas compositions for the CH$_4$ and H$_2$O formation. Furthermore, two adsorbed intermediate species with different reactivity are postulated for CO containing feeds, while only one is present for CO$_2$ methanation.

Hence, the results obtained via the PTK method provide the experimental basis to identify, parameterize and validate the surface reaction steps being significant for an appropriate transient kinetic model. The present work focuses on the impact of the CO/CO$_2$ ratio at a constant CO$_x$/H$_2$ ratio on the transient response, in order to demonstrate the capabilities of the PTK method to elucidate mechanistic aspects. For deeper investigation of the dynamic processes on the catalyst surface, however, the variation of the CO$_x$/H$_2$ ratio and the pressure is necessary, as well.

The reported results are based on observations at the macroscopic level by analyzing the transient molar flow rates at the reactor outlet. Therefore, complementary *in situ* or *operando* measurements with emphasis on the dynamics of processes at the solid surfaces are required for a full kinetic understanding of the dynamic reactor behavior, which are matter of ongoing research. This knowledge provides the basis to derive reaction kinetics valid under unsteady-state conditions mandatory to predict the behavior of methanation reactors in PtG application scenarios. In particular, a micro-kinetic model is recommended, which also considers the storage capacity of the reactants at the active material and support. The model improved by appropriate micro-kinetics also allows the design of suitable reactors offering sufficient tolerance towards fluctuating feed gas compositions. In principle, the reported methodology can also be applied to other heterogeneously catalyzed reactions directly, e.g. ammonia or methanol synthesis, also relevant for chemical storage of renewable energy.

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