On the kinetics and equilibria of plasma-based dry reforming of methane

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HIGHLIGHTS

- CO\textsubscript{2} and CH\textsubscript{4} reforming in plasma is characterized in terms of equilibrium and kinetics.
- CO\textsubscript{2} splitting has favourable kinetics while CH\textsubscript{4} reforming exhibits high conversions.
- Combining the gasses in DRM shows a synergy leading to optimal rates and conversions.
- Addition of a packing material has gas mixture-specific effects to tune the kinetics.
- The DRM gas mixture ratio is key to tune the kinetics and product distribution.

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ABSTRACT

Plasma reactors are interesting for gas-based chemical conversion but the fundamental relation between the plasma chemistry and selected conditions remains poorly understood. Apparent kinetic parameters for the loss and formation processes of individual components of gas conversion processes, can however be extracted by performing experiments in an extended residence time range (2–75 s) and fitting the gas composition to a first-order kinetic model of the evolution towards partial chemical equilibrium (PCE). We specifically investigated the differences in kinetic characteristics and PCE state of the CO\textsubscript{2} dissociation and CH\textsubscript{4} reforming reactions in a dielectric barrier discharge reactor (DBD), how these are mutually affected when combining both gases in the dry reforming of methane (DRM) reaction, and how they change when a packing material (non-porous SiO\textsubscript{2}) is added to the reactor. We find that CO\textsubscript{2} dissociation is characterized by a comparatively high reaction rate of 0.120 s\textsuperscript{-1} compared to CH\textsubscript{4} reforming at 0.041 s\textsuperscript{-1}; whereas CH\textsubscript{4} reforming reaches higher equilibrium conversions, 82\% compared to 53.6\% for CO\textsubscript{2} dissociation. Combining both feed gases makes the DRM reaction to proceed at a relatively high rate (0.088 s\textsuperscript{-1}) and high conversion (75.4\%) compared to CO\textsubscript{2} dissociation, through accessing new chemical pathways between the products of CO\textsubscript{2} and CH\textsubscript{4}. The addition of the packing material can also distinctly influence the conversion rate and position of the equilibrium, but its precise effect depends strongly on the gas composition. Comparing different CO\textsubscript{2}:CH\textsubscript{4} ratios reveals the delicate balance of the combined chemistry. CO\textsubscript{2} drives the loss reactions in DRM, whereas CH\textsubscript{4} in the mixture suppresses back reactions. As a result, our methodology provides some of the insight necessary to systematically tune the conversion process.

1. Introduction

Plasmas offer an interesting approach to gas conversion for industrial applications [1-3]. Their highly energetic, collision-rich environment is often employed to activate highly stable compounds through (strongly endothermic) reactions, such as CO\textsubscript{2} splitting [4-10], dry reforming of methane (DRM) [4,11-13], and nitrogen fixation [14-16], at milder and more convenient conditions (ambient pressure and...
temperature) than for thermal conversion. Multiple reactor types have already been investigated, e.g. microwave discharge [17,18], dielectric barrier discharge (DBD) [5-12], and gliding arc discharge [19,20]. Altering the specific design of the DBD reactor [7,21-24], changing multiple process parameters [25], and adding (catalytic) packing materials [6,7,26-28] have all shown interesting and promising results. However, the fundamental relation between the plasma chemistry and the selected plasma conditions and reactor parameters remains poorly understood.

Many of these previous studies claimed that plasma processes may enable thermodynamically unfavourable conversions beyond the thermal limit and at higher rates than conventional thermal approaches. In spite of these promising observations, however, further optimization of plasma-based conversion processes is very difficult because their most fundamental characteristics are unknown. For thermal processes key quantities such as equilibrium constants or reaction rates of the main reaction channels are generally known and can be used to predict the yield in a given process setup. They are, however, usually not measured for plasma-based conversion processes. Most of the time, parametric studies of plasma-based gas conversions only report conversions or gas compositions at one or a limited set of residence times, which precludes elucidation of the kinetics or maximal yield, and makes it difficult to conclusively identify synergies between plasma and (catalytic) packing materials.

Despite the intrinsically nonequilibrium nature of plasmas, it is however possible to characterize a plasma-based conversion process in terms of a so-called partial chemical equilibrium (PCE) without the need to know the exact reaction schemes, as we have recently demonstrated for CO2 dissociation in a DBD reactor [25]. Here, we found that the concentration of CO2, CO, and O2 in the driven, out-of-equilibrium plasma evolves to a final composition in a similar manner as a thermal system would evolve to its equilibrium state, i.e. the same final composition is reached when starting from a pure CO2 flow or a stoichiometric mixture of CO and O2. The position of the PCE is specific to the discharge conditions at hand, as was also suggested by Vepřek and co-workers [29-31]. Note that in all cases the measured PCE state corresponded to a large CO2 conversion (up to 70% in a SiO2-packed 455 µm reactor [25]), which would thermally only be attainable at very high temperatures (>3000 K) [4]. A simple PCE-based approach can show that the kinetics of plasma (catalysis) are different from traditional thermal kinetics, but can still be described by the same key criteria (i.e. rate coefficient and equilibrium). In contrast to thermal processes, these are however affected by more and different parameters, allowing for additional flexibility to tune the maximal conversion and rate.

The interesting observations of [25] can form the basis for more elaborate research into more complex chemistry sets, such as dry reforming of methane (DRM). Indeed, CH4 reforming on its own already shows a more complex chemistry, compared to pure CO2 conversion, since its products vary from C to H2 to CxHy molecules, with thousands of reactions among them [32]. High conversions of methane up to 80% have been reported, with hydrocarbons varying results [11,12,20,23,37-39]. However, the highly complex chemistry of plasma-based DRM, combined with its technological potential, require a deeper insight into the fundamental characteristics of the process. A lot of knowledge is available from detailed reaction schemes obtained through modelling, e.g. those in [32,40,41], but they only partially capture the complexity of experimental reactors and how set-up parameters change the overall kinetics.

In this work we will therefore investigate how the kinetics of CO2 dissociation change upon addition of CH4 in a DBD reactor. For this purpose, we generalise our PCE model (that was specifically developed and only applicable for CO2 splitting) to arbitrary gas mixtures with a priori unknown stoichiometry. Hence, this novel method does not require knowledge on the precise process stoichiometry, and can be applied to multicomponent mixtures, also for other gas conversion applications. This approach will allow other researchers as well to compare the conversion characteristics of certain individual components across different reactors and gas mixtures.

In order to understand the occurring changes, pure CH4 reforming will be tested and compared at the same conditions as used in our previous work on CO2 splitting [25] being the benchmark. In this way we can elucidate how the apparent kinetics of these pure reactions are different from their combination in the DRM reaction. Also, we will evaluate the impact of a non-porous SiO2 spheres as packing on the kinetics of CH4 reforming and DRM, compared with CO2 dissociation, and how the CO2:CH4 ratio in DRM shifts the kinetics. Despite the simplicity of the experimental set-up, salient mechanistic insights are obtained. Eventually, information on the maximal conversion in a DBD plasma and its overall kinetics will be useful in the determination of optimal process conditions, and to compare different set-ups and packing materials on a systematic basis.

2. Methods and theory

2.1. Experimental set-up

The experiments were performed in the same set-up as in our previous work [25], as shown in Fig. 1. A 455 µm gap size was used as the reference gap size, also showing the highest rate coefficients of the various gap sizes tested. We used a coaxial DBD micro gas reactor, made of an alumina dielectric tube with a precision honed inner diameter of 17.41 mm and a wall thickness of 2.5 mm. A grounded stainless-steel rod with an outer diameter of 16.50 mm was placed in the centre of the alumina tube, to act as the inner electrode and to shape the reaction volume with a discharge gap of 455 µm. The tube was wrapped with a stainless-steel mesh over a length of 100 mm to act as the high voltage outer electrode of the DBD reactor and define the reaction zone length. The resulting reaction volume is 2.4 × 10^5 mm^3. Besides studying the empty reactor, the reactor was also packed with non-porous SiO2 spheres (Sigmund Lindner) with a size range of 100–200 µm. We used a SiO2 packing to be able to compare with our previous results for pure CO2 splitting, and because the other packing material used in our previous work (ZrO2) did not reach the desired 30 W (triggering the current safety switch of the power supply) in CH4 or CO2/CH4 mixtures for DRM. Although many different materials have been studied in literature in packed bed DBD, we focus on SiO2. Indeed, the inert nature of non-porous SiO2 is beneficial to minimise any additional effects that a packing material could have on the reactor performance (e.g. catalysis), and to focus on the basic kinetics for this study. The packing was added to the reactor and vibrated for one minute via an external device to ensure the closest packing possible in a repeatable fashion. No significant material degradation was observed during the experiments.

A high voltage was applied by generating a 3 kHz sine wave by a PC-controlled function generator (Tektronix, AFG 2021) and amplifying the signal with a high voltage amplifier (TREK, Model 20/20C-HS, x2000 voltage amplification). Continuous power measurements, recorded with a digital oscilloscope (Picotech, Picoscope 6402D2), a high voltage probe (Tektronix, P6015A), and a current transformer (Pearson, Model 4100), were used to adjust the sine wave amplitude (between 15 and 20 kV) to obtain and maintain the desired power of 30 W. This power was measured during a number (n) of consecutive periods (T):

$$P = \frac{1}{RT} \int_{0}^{\infty} U(t)I(t)dt$$

(1)
The reactor was fed with pure CO$_2$, CH$_4$, or a mixture of both, at different flow rates to obtain the desired residence times between 2 and 75 s in the reactor. The flow rates were set and monitored by two mass flow controllers (Bronkhorst EL-FLOW Select series). The pressure in the tubing between the reactor and the GC was kept at 1.2 bara ± 0.2 bara. In the case of the SiO$_2$ packed reactor, adjusted, lower flow rates were used to account for the loss in reaction volume by the packing. A packing efficiency of 49.5% was estimated, based on the computational results of our previous work for the identical case of 100–200 µm spheres in the 455 µm gap [25]. Note that we have no data point at 70 s for the SiO$_2$ packed DRM experiment. This point could not be measured due to the lower limit of the MFC, being 0.65 mL/min. Note that alteration of the flow rate was done to achieve different residence times, rather than changing the reactor length, in order to maintain the same power density, i.e. power-to-volume ratio, and thus constant plasma characteristics. While changing the flow rate might affect the mass and heat transfer rates, the gas temperature was estimated only slightly above room temperature (based on the measured temperature immediately after shutdown, being always below 50 °C), so heat transfer should not be an issue here, like it could be in thermal reactors.

The gaseous products were analysed by a gas chromatograph (Compact GC, Interscience) with pressure-less sampling. This GC has two thermal conductivity detector (TCD) channels and one flame ionization detector (FID). The first TCD channel (TCD B) contains a Rt-Q-Bond column able to separate CO$_2$ and large hydrocarbons from permanent gases, the second TCD channel (TCD M) contains a Rt-Q-bond pre-column to delay CO$_2$ and the larger components and only inject the permanent gases on a Molsieve 5A column and separate them. The FID has a Rt-x-1, 5 µm column to separate and detect (oxygenated) hydrocarbons. CO$_2$, CO, O$_2$, CH$_4$, H$_2$, C$_2$H$_6$, C$_3$H$_8$, C$_2$H$_5$OH, and C$_2$H$_4$OH were calibrated by using calibration standards (Air Liquide). More peaks could be identified but were not calibrated, since either no calibration standard was available (vapor injection identification) or they coincide with other peaks; they will not be discussed in the main text but the results are available in section 2 and 3 of the supplementary material. The CO$_2$ or CH$_4$ conversion derived from the GC data was defined as:

$$X = \frac{\dot{y}_i - \dot{y}_{ini}}{\dot{y}_{ini}}$$

with $\dot{y}$ the molar flow rate of component $y$, being either CO$_2$ or CH$_4$. The total conversion was calculated according to the CO$_2$:CH$_4$ molar ratio of the mixture, A:B, as:

$$X_{total} = \frac{AX_{CO_2} + BX_{CH_4}}{A + B}$$

2.2. Experimental method

The standard experimental conditions in this work are set at 30 W and 3 kHz, a 455 µm discharge gap size, and 1.2 bara, at various residence times up to 75 s. The reactor was operated for at least 40 min to let it reach a thermal steady-state behaviour, i.e. a stable reactor temperature and voltage is reached due to heat losses towards the reactor and environment. Extended operating times up to 120 min were used for flow rates below 10 mL/min, to ensure steady-state behaviour in the reactor and following tubing, for consistent gas composition analysis. The applied voltage was periodically adjusted on the function generator to obtain and maintain the desired constant plasma power of 30 W. Four GC and oscilloscope measurements were recorded as soon as the steady-state behaviour from above was reached to evaluate the GC error.

Every condition was tested in three-fold for statistical review of the error, due to packing of the reactor, plasma stabilization, and random (environmental) effects. The error bars are defined as:

$$error = \pm S_n \frac{T_{t}(p, n_s)}{\sqrt{n_i}}$$

with $S_n$ the sample standard deviation of the measurements, $n_s$ the sample size, and $T_{t}$ the two-tailed inverse of the Student’s t-distribution for sample size $n_i$ and probability $p$ set at 95%. 2.3. Partial chemical equilibrium (PCE)

The overall reaction rate coefficient and location of the PCE are determined by applying an apparent first order reversible reaction fit to the residence time measurements. In our previous work, we explicitly derived an expression for the time evolution of molecular concentrations towards the PCE state of the CO$_2$/CO/O$_2$ system, wherein we assumed first order kinetics and stoichiometric conversion between the aforementioned molecules only [25]. Such an approach is no longer valid for more complex gas mixtures, such as those obtained for CH$_4$...
conversion or DRM. However, a slight generalization of the equilibrium model permits its application to the conversion of any molecule in an arbitrary gas mixture, without requiring any detailed information on reaction products or mechanisms. Only first order kinetics—reaction rate proportional to the concentration of reacting molecules—is assumed. Each process could progress through many different possible individual mechanisms, between which we cannot distinguish experimentally. The measured rate coefficient is therefore a weighted average for all these individual reactions which, by construction, is assumed to be constant over time.

The loss and formation rate of a gas molecule $A$ through unspecified pathways with rate coefficients $k_{\text{loss}}$ and $k_{\text{form}}$ are:

$$r_{A\text{,loss}}(x_A) = k_{\text{loss}}x_A$$

(5)

$$r_{A\text{,form}}(x_A) = k_{\text{form}}(1 - x_A)f$$

(6)

Not all of the (non-$A$) molecules in the system can directly be converted into $A$, because only a fraction $f$ has the “right” reactivity. We assume that $f$ is a constant that depends on the elemental composition of the system. By construction, it is therefore a time-averaged stoichiometric parameter throughout all stages of the conversion process, just like the rate coefficient to which it is tied. Derivation of the formula, that is shown in detail in the supplementary information leads to the fit equation describing the concentration (in mole fraction) of $A$ in time:

$$x_A(t) = x_{A,e} - (x_{A,e} - x_A)e^{-kt}$$

(7)

where $x_{A,e}$ and $x_{A,i}$ are the equilibrium and initial mole fraction of $A$ and $k = f k_{\text{form}} + k_{\text{loss}}$.

(8)

This equilibrium mole fraction $x_{A,e}$ can be rewritten in terms of the total equilibrium conversion $X_{\text{Total,e}}$ as:

$$X_{\text{Total,e}} = \frac{k_{\text{loss}}}{f k_{\text{form}} + k_{\text{loss}}}$$

(9)

The fit was applied by importing the experimental data (consisting of up to 132 data points per parameter and reaction) into MATLAB as gas fractions, calculating a fit according to Eq. (7) resulting in $k$ and $x_{A,e}$, and finally converted back into conversions for plotting on the graphs. $x_{A,i}$ is equal to 1 for pure CO2 and CH4 as well as for fitting the overall DRM trend (where $x_{A} = x_{\text{CO2}} + x_{\text{CH4}}$), while the respective CO2 and CH4 fractions were taken for their individual fits in mixed DRM cases. $k_{\text{loss}}$ and $f k_{\text{form}}$ are calculated from Eqs. (8) and (9).

3. Results and discussion

3.1. The benchmark: Separate CO2 and CH4 conversion

CO2 dissociation was performed in our previous work [25] at the standard conditions mentioned in the experimental method. Note that the thermodynamic equilibrium at the considered (ambient) temperature and pressure is virtually 0% for all gas compositions (pure CO2 and CH4, as well as all DRM mixtures) [4]. The results, plotted as the black curve in Fig. 2(a), however show that the CO2 splitting reaction under the considered plasma conditions was characterized by an equilibrium conversion of 53.6% with an apparent overall reaction rate coefficient of 0.120 s$^{-1}$ (Table 1), found by the apparent first order reversible reaction fit. Viewed through the lens of our simplified equilibrium model, and more specifically the expressions for the apparent reaction rate coefficient and equilibrium conversion in equations (8) and (9), we get a $k_{\text{loss}}$ term of 0.066 s$^{-1}$ and a $f k_{\text{form}}$ term of 0.056 s$^{-1}$. The dominant reactions in the DBD reactor are electron-impact dissociation of CO2 as main CO2 loss process, while three-body recombination of CO and O is the most important formation reaction, as reported by Aerts et al. [8].

CH4 reforming was carried out at the same conditions, and the results are shown in Fig. 2(a) and Table 1 as well. They exhibit completely different behaviour from CO2 dissociation with a higher equilibrium conversion of 82%. However, the overall apparent reaction rate coefficient of CH4 reforming is only a third of the value for CO2 dissociation, i.e. 0.041 s$^{-1}$. Our equilibrium model shows that $k_{\text{loss}}$ and $f k_{\text{form}}$ both have much lower values than those seen with CO2 dissociation, being 0.034 s$^{-1}$ and 0.0073 s$^{-1}$ respectively (cf. equations (8) and (9)). From our modelling work it was established that the CH4 loss reactions in a DBD reactor are dominated by electron impact ionization and dissociation of CH4, yielding CH4$^+$ and CH3$^+$ ions, as well as CH3, CH2, and CH radicals, cfr. equations R11-5 in [32] and Table 2 in [41]. In view of the small value of $k_{\text{loss}}$, these processes are therefore not as efficient as in the pure CO2 plasma, which seems counterintuitive based on the reaction enthalpies. However, as shown by Snoeckx et al. [41], electron impact dissociation of CH4 is very fast, but the recombination of the produced CH4 radicals back towards CH4 is also very quick, resulting in a net low conversion and thus low effective rate coefficient. On the other hand, the products of CO2 electron impact dissociation are more stable: CO is a saturated molecule, and the O atoms can recombine into O2, ultimately leading to the higher effective loss rate from electron impact-driven processes.

The net CH4 formation is mainly due to electron impact dissociation of C2H2 and C2H4 (cfr. equations R12 and R14 in [32]). These C3 hydrocarbons are however not very abundant in our CH4 plasma, with a maximum product fraction of 2.46% measured in all our CH4 experiments at steady state (see Table S 3 in the supplementary material). In addition, experiments in the packed reactor (section 3.3) indeed confirm that both loss and formation processes are primarily electron impact-driven. Essentially, this means that CH4 formation is hampered by a small $f$ factor, which leads to a smaller effective formation rate coefficient $f k_{\text{form}}$, thus shifting the conversion equilibrium away from CH4. Because the electron impact-driven processes in this system are not as efficient as in the CO2 system, the overall evolution rate towards this high equilibrium conversion is however lower.

Note that, although the plasma chemistry processes on the macroscale are inherently fast in nature (milliseconds), the conversion on the macroscale is still a rather slow process in this DBD reactor. This is due to the balance act of short and select random micro-discharges per second (with a diameter of typically 100 µm and duration of about 200 ns), in combination with reactions in the afterglow, diffusion, convection, and back reactions, resulting an overall slower conversion process, requiring about 40 s to 80 s to reach PCE, depending on the composition.

3.2. DRM: The best of both worlds

Subsequently, both gases were combined in an equimolar ratio to perform the DRM reaction at the same standard conditions (Fig. 2(a) and Table 1). The combination of CO2 and CH4 leads to a high overall equilibrium conversion of 75.4%, associated with a high rate coefficient of 0.088 s$^{-1}$, which is higher than the numerical average of the individual reactions would have been. This is a manifestation of $k_{\text{loss}}$ being high at 0.066 s$^{-1}$, which is the same as for CO2 dissociation, and $f k_{\text{form}}$ staying fairly low at 0.0217 s$^{-1}$. When performing a similar analysis purely on the respective concentrations of CO2 and CH4 in DRM, we can see that, compared to the pure gases, $k_{\text{loss,CO2}}$, $k_{\text{loss,CH4}}$, $f k_{\text{form,CO2}}$, and $f k_{\text{form,CH4}}$ are more than two times larger, whereas $k_{\text{form,CO2}}$ is more than halved. This means that additional reaction pathways have become available for the processes producing and consuming CH4, while CO2 formation pathways are suppressed due to the mixing of the two gases. These observations were also predicted by kinetic modelling [41,42]. The density of O atoms is very low in the CO2:CH4 mixture (there is also less than 0.023% O2, see section 3.5.3) and this limits CO2 formation by
lowering its $f$ factor, an effect that cannot be fully compensated by new “combustion-type” reactions of CH4. These latter reactions, however, do affect the consumption of CH4 because it can react with species such as O, OH and CO2. CH4 formation benefits from the availability of more radicals for three-body recombinations in DRM [41], hence explaining the increase of $f_{\text{CH4},\text{form,CH4}}$ when CO2 is present.

These changes result in mixing of the characteristics of the individual reactions, towards the overall DRM kinetics trend seen above. In total, we effectively see CO2 boosting the conversion rate coefficient of CH4 while it reduces its own rate coefficient although to a smaller extent. Moreover, CH4 addition increases the (equilibrium) conversion of CO2 while slightly reducing its own equilibrium conversion. The overall result is that we obtain the best of both worlds in DRM, i.e. high apparent rate coefficients, like in CO2 dissociation, and high equilibrium conversions, like in CH4 reforming. In the most practical sense, the changes in kinetics and PCE state, resulting from mixing CO2 and CH4, ensure that for any “practical” (short) residence time, the DRM process significantly outperforms its pure gas counterparts in terms of obtained conversion for both compounds, as can be seen in Fig. 2(a).

It should also be noted that the individual CO2 and CH4 conversion in our DRM reaction, under the applied conditions here, are very close to each other, in comparison to other studies [11,12,43,44]. These papers report typical $X_{\text{CH4}}:X_{\text{CO2}}$ ratios between 1.5 and 2, while ours are between 1.22 and 1.12 at the shortest and longest residence time, respectively.

### Table 1

Fitted kinetic and partial chemical equilibrium data for CO2 dissociation, CH4 reforming, and DRM, as well as its sub-reactions, for both the empty and SiO2-packed reactor at the standard conditions.

<table>
<thead>
<tr>
<th>Reaction</th>
<th>CO2 splitting [25]</th>
<th>CH4 reforming</th>
<th>DRM total</th>
<th>DRM CO2</th>
<th>DRM CH4</th>
</tr>
</thead>
<tbody>
<tr>
<td>$k$ (s⁻¹)</td>
<td>0.120 ± 0.005</td>
<td>0.041 ± 0.002</td>
<td>0.088 ± 0.003</td>
<td>0.083 ± 0.002</td>
<td>0.093 ± 0.003</td>
</tr>
<tr>
<td>$k_{\text{CO2}}$ (s⁻¹)</td>
<td>0.066 ± 0.003</td>
<td>0.034 ± 0.003</td>
<td>0.066 ± 0.003</td>
<td>0.059 ± 0.003</td>
<td>0.074 ± 0.003</td>
</tr>
<tr>
<td>$f_{\text{form}}$ (s⁻¹)</td>
<td>0.056 ± 0.003</td>
<td>0.0073 ± 0.0006</td>
<td>0.0217 ± 0.0009</td>
<td>0.024 ± 0.001</td>
<td>0.0190 ± 0.0008</td>
</tr>
<tr>
<td>$X_e$ (%)</td>
<td>53.6 ± 0.8</td>
<td>82 ± 2</td>
<td>75.4 ± 0.6</td>
<td>71.2 ± 0.7</td>
<td>79.7 ± 0.7</td>
</tr>
</tbody>
</table>

### 3.3. Tuning the kinetics by packing material

Our previous work showed that adding a packing material to the reactor can tune both the rate coefficient and equilibrium conversion of plasma-induced CO2 dissociation individually, while at most enhancing only the rate in traditional thermal reactors (in case of a catalytic packing) [25]. These changes are usually attributed to alteration of the plasma properties by physical effects of the packing materials on the discharge [45]: When a plasma changes, also the associated PCE can change. Generally speaking, adding a packing material will increase the local electric field near the contact points, and thus increase the electron temperature, while slightly lowering densities of some species due to more surface losses, depending on the material-gap combination [46]. Indeed, the applied peak-to-peak voltage (a measure for the reduced electric field E/N as the discharge gap remains constant and thus also for the electron temperature) increases significantly for all compositions upon adding the SiO2 packing, see Table S 10 in the supplementary information. This should result in a boost of the electron impact reactions, i.e. the major loss reactions for CO2 and CH4, therefore increasing $k_{\text{CO2}}$, $k$, and $X_e$.

A SiO2 packing was previously found to significantly increase the CO2 equilibrium conversion from 53.6% to 71.1%, while only slightly decreasing the rate coefficient from 0.120 s⁻¹ to 0.111 s⁻¹ [25], as shown when comparing Fig. 2 (a) and (b) and Table 1. Indeed, $k_{\text{CO2}}$ increases because electron impact-based loss reactions are stimulated by the increased electron temperature, whereas $f_{\text{form}}$ does not, because three-body neutral recombinations are unaffected by the electron temperature [8]. In fact, $f_{\text{form}}$ even decreases, which may be explained...
by the larger surface/volume ratio in the packed reactor: the presence of a large surface area will facilitate surface-mediated O atom recombination, thus reducing the density of O atoms in the gas phase, and hence suppressing the formation of CO\textsubscript{2} along the three-body recombination pathway by lowering the $f$ factor \cite{47}. The opposite effect of the packing on the respective CO\textsubscript{2} dissociation and formation rates therefore explains how the CO\textsubscript{2} equilibrium conversion can increase, while its overall conversion rate coefficient slightly decreases.

The same experiments for CH\textsubscript{4} reforming show a (almost) doubling of both $k_{\text{loss}}$ and $f_{\text{form}}$ upon packing the reactor. While the SiO\textsubscript{2} packing does not enhance the equilibrium conversion, the apparent rate coefficient almost doubles from 0.041 s\textsuperscript{-1} to 0.074 s\textsuperscript{-1} (see Table 1). As discussed in section 3.1, the most prominent reactions for both loss and formation of CH\textsubscript{4} in a DBD are again electron impact reactions—i.e. dissociation and ionization for CH\textsubscript{4} loss and dissociation of C\textsubscript{3}H\textsubscript{6} and C\textsubscript{3}H\textsubscript{8} for CH\textsubscript{4} formation, respectively. This means that both loss and formation are equally affected by a change in electron temperature, so that the ratio of $k_{\text{loss}}$ and $f_{\text{form}}$ is constant (as well as the equilibrium conversion), but their sum (and the overall reaction rate) increases.

Finally, the addition of a SiO\textsubscript{2} packing in DRM increases both the total equilibrium conversion, from 75.4\% to 84.3\%, and the apparent total rate coefficient, from 0.088 s\textsuperscript{-1} to 0.130 s\textsuperscript{-1} (cf. Table 1). Although the individual CO\textsubscript{2} and CH\textsubscript{4} rate coefficients (0.118 s\textsuperscript{-1} and 0.143 s\textsuperscript{-1} respectively) differ more compared to the empty reactor (section 3.2), the individual equilibrium conversions become roughly equal (84.3\%), which is different from the empty reactor where the CH\textsubscript{4} conversion remained higher that the CO\textsubscript{2} conversion. This results in $X_{\text{CH}_4}:X_{\text{CO}_2}$ ratios being even closer to 1, i.e. 1.29 to 0.99 at the shortest and longest residence times, respectively. The data shows that for both CO\textsubscript{2} and CH\textsubscript{4}, and thus in total for DRM as well, $k_{\text{loss}}$ strongly increases while $f_{\text{form}}$ stays about the same when the packing material is added. The changes in kinetics are again expected since both CO\textsubscript{2} and CH\textsubscript{4} losses are based on electron impact reactions, resulting in a rise of $k_{\text{loss}}$ due to the higher electron temperature in case of the SiO\textsubscript{2} packing. The formation reactions have been predicted to be mostly based on ion and neutral chemistry for CO\textsubscript{2}, and three-body recombination, electron impact on C3 (which are barely present), and C2 three-body reactions for CH\textsubscript{4}, meaning that $f_{\text{form}}$ is almost unaffected \cite{41,42}.

### Table 2

<table>
<thead>
<tr>
<th>CO\textsubscript{2} : CH\textsubscript{4} ratio</th>
<th>1:0 [25]</th>
<th>3:1</th>
<th>1:1</th>
<th>1:3</th>
<th>0:1</th>
</tr>
</thead>
<tbody>
<tr>
<td>$k$ (s$^{-1}$)</td>
<td>0.120 ± 0.005</td>
<td>0.108 ± 0.004</td>
<td>0.088 ± 0.003</td>
<td>0.080 ± 0.002</td>
<td>0.041 ± 0.002</td>
</tr>
<tr>
<td>$k_{\text{loss}}$ (s$^{-1}$)</td>
<td>0.066 ± 0.003</td>
<td>0.065 ± 0.003</td>
<td>0.066 ± 0.003</td>
<td>0.066 ± 0.002</td>
<td>0.034 ± 0.003</td>
</tr>
<tr>
<td>$f_{\text{form}}$ (s$^{-1}$)</td>
<td>0.056 ± 0.003</td>
<td>0.042 ± 0.002</td>
<td>0.0217 ± 0.0009</td>
<td>0.0136 ± 0.0005</td>
<td>0.0073 ± 0.0006</td>
</tr>
<tr>
<td>$X_{\text{C}_2\text{H}_4}$ (%)</td>
<td>53.6 ± 0.8</td>
<td>60.5 ± 0.6</td>
<td>75.4 ± 0.6</td>
<td>83.0 ± 0.7</td>
<td>82 ± 2</td>
</tr>
</tbody>
</table>

### 3.4. Tuning the kinetics by mixing ratio

In section 3.1 we discussed how CO\textsubscript{2} dissociation is characterized by a high apparent rate coefficient but low equilibrium conversion, while CH\textsubscript{4} reforming displays a low apparent rate coefficient but high equilibrium conversion. Combining both in DRM results in total rates and conversions in between these values. In this section we look further into the mechanistic aspects of this mixing, by also testing 3:1 and 1:3 CO\textsubscript{2}:CH\textsubscript{4} ratios.

The time evolution of the conversions shown in Fig. 3—with their associated kinetic and equilibrium parameters compared to the pure gases in Table 2—further clarifies the extent of the mixing effect seen in section 3.2. The rate constants and equilibrium conversion for the different gas mixing ratios interpolate continuously, but not linearly, between those for pure CO\textsubscript{2} and pure CH\textsubscript{4}. The more CO\textsubscript{2} is added to the
mixture, the higher the overall rate coefficient is, skewed towards that of the pure CO2 system. The equilibrium conversion, however, is mainly affected by the CH4 concentration (i.e. skewed towards the pure CH4 case), with higher equilibrium conversions obtained with more CH4 in the mixture. The changes in kloss and f\textsubscript{f,prem} curiously show that as long as there is CO\textsubscript{2} in the mixture, the loss rate coefficient kloss of DRM is constant and equal to kloss of pure CO\textsubscript{2}. At first sight, the shifts in k and X\textsubscript{prem} as a function of the CO\textsubscript{2}-CH\textsubscript{4} ratio may therefore seem to be fully attributable to a change of f\textsubscript{f,prem} which decreases with increasing CH\textsubscript{4} concentration. Analysis of the effective kinetics of CO\textsubscript{2} and CH\textsubscript{4} shown in Table 2 (based on their individual conversion within DRM), however, shows how these shifts of k and X\textsubscript{prem} are skewed, and those will be discussed here. The results of the different gas mixing ratios investigated. Because the CH\textsubscript{4} loss is enhanced by species produced in CO\textsubscript{2} loss, concentration, although at the 3:1 ratio, f\textsubscript{f,prem,CH\textsubscript{4}} at the 3:1 ratio is close to the value for pure CO\textsubscript{2}. As a result, the 3:1 mixture exhibits a much higher conversion of CH\textsubscript{4} than of CO\textsubscript{2} (89.0% and 51.1%, respectively). The formation rate coefficients for both gases generally decrease with increasing CH\textsubscript{4} concentration, although at the 3:1 ratio, f\textsubscript{f,prem,CO\textsubscript{2}} is lower than at the 1:1 ratio, whereas f\textsubscript{f,CO2,CH\textsubscript{4}} at the 3:1 ratio is close to the value for pure CO\textsubscript{2}. As a result, the 3:1 mixture exhibits a much higher conversion of CH\textsubscript{4} than of CO\textsubscript{2} (89.0% and 51.1%, respectively).

In the most general sense, changing the CO\textsubscript{2}-CH\textsubscript{4} ratio therefore mainly changes the formation aspect of the overall reaction, allowing us to tune both the kinetics as well as the PCE of DRM. From the trends in the loss and formation rates, we conclude that CO\textsubscript{2} drives the loss reactions in DRM, whereas the presence of CH\textsubscript{4} in the mixture suppresses formation reactions. The combination of these effects results in both high equilibrium conversions and high conversion rates in DRM. Combining this knowledge with insight gleaned from the experiments with the SiO\textsubscript{2}-packed reactor, it can be inferred that electron impact-driven processes are very efficient with CO\textsubscript{2} and they dominate the non-equilibrium loss processes even at fairly low CO\textsubscript{2} fractions (25% in our experiments). CH\textsubscript{4} loss is enhanced by species produced in CO\textsubscript{2} loss reactions which, in turn, chemically suppress CO\textsubscript{2} formation. To have an appreciable effect on the CO\textsubscript{2} conversion, the CH\textsubscript{4} fraction must however be sufficiently high (50% in our experiments).

Table S 10 in the supplementary material shows the applied peak-to-peak voltage for the different gas mixing ratios investigated. Because we keep the discharge gap constant, the peak-to-peak voltage can be considered as a measure for the reduced electric field, and thus for the electron temperature. In contrast to the addition of the SiO\textsubscript{2} packing, we see no clear correlation between this peak-to-peak voltage and the kinetic data derived in our study, for the different gas mixtures. This indicates that besides the electron temperature, additional unknown factors will have an impact as well.

3.5. How residence time and gas mixture tune product composition

Besides the CO\textsubscript{2}, CH\textsubscript{4}, and total conversions reported above, it is interesting to understand how these differences influence the formed products. As mentioned in section 2.1, only a number of components could be quantified, and those will be discussed here. The results of the remaining components can be found with their peak areas in the supplementary material. During the experiments with CH\textsubscript{4}, mostly at longer residence times, we could collect small amounts of carbon deposition and highly viscous fluids. GC-MS analysis showed ppm levels of C1 to C4 acids and unidentified carbon polymers. This and the uncalibrated components will of course result in an incomplete atom balance after the reactor, which is further discussed in section 3.6.1.

3.5.1. CO\textsubscript{2} dissociation

The products formed in CO\textsubscript{2} dissociation are limited to CO, O\textsubscript{2}, O\textsubscript{3}, and C deposits. Only trace amounts of O\textsubscript{3} were detected and no C deposition was observed in or after the reactor, so these will be neglected further on. The production of CO and O\textsubscript{2} in the empty reactor is shown in Fig. 4(a). We can see that the production of CO and O\textsubscript{2} respects the stoichiometric 2:1 ratio of the CO\textsubscript{2} splitting reaction, with CO reaching a maximum concentration of around 36% and O\textsubscript{2} reaching around 18%.

Adding a SiO\textsubscript{2} packing to the reactor enhances the conversion, as discussed in section 3.3, which also results in higher CO and O\textsubscript{2} concentrations, as shown in Fig. 4(b). The maximum CO concentration obtained, increases to about 49% while the O\textsubscript{2} concentration increases to 24%, again respecting the stoichiometric 2:1 ratio of the CO\textsubscript{2} splitting reaction.

3.5.2. CH\textsubscript{4} reforming

Fig. 4(c) shows the calibrated product composition after CH\textsubscript{4} reforming for the empty reactor (the exact concentration of every component can again be found in Table S 3 in the supplementary material). The main component formed during CH\textsubscript{4} reforming is H\textsubscript{2}, followed by ethane, propane, and ethene. A steady increase in H\textsubscript{2} concentration is seen from 8% to 74% upon increasing the residence time, following the increasing conversion. This trend is, however, not followed by the other components, of which the concentration first increases to a maximum value, and then decreases again. Ethene reaches a maximum concentration of 6.21% at 40 s, and slightly decreases to 5.5% at 70 s. Propane shows the same behaviour with a maximum concentration of 2.46% at 40 s, dropping to 2.1% at 70 s. Ethane reaches its maximum concentration of 0.79% already at 17.5 s, and decreases significantly to only 0.35% at 70 s. This means that these small hydrocarbons are actually intermediates that are consumed, either in the production of higher hydrocarbons or back into formation of CH\textsubscript{4}, as well as H\textsubscript{2} and C, at longer residence times. Also, it can be inferred that alkenes are more prone to react further or be decomposed again, since they are more reactive.

Adding the SiO\textsubscript{2} packing to the reactor zone has two major effects on the product composition after CH\textsubscript{4} reforming. A higher share of H\textsubscript{2} is observed over the entire residence time range, but the changes in C2 and C3 concentration depend on the exact residence time and there are also lower amounts of all higher hydrocarbons.

3.5.3. Dry reforming of methane

Fig. 5 shows the calibrated product composition for DRM. All concentrations can again be found in the supplementary material (Table S 6). The addition of CO\textsubscript{2} to the mixture results in high fractions of CO in the reactor effluent, as it is the main product from electron impact dissociation of CO\textsubscript{2}. The concentration starts at 10.2% at the shortest residence time and rises to 37.4% at the longest residence time. The other main product from CO\textsubscript{2} dissociation, O\textsubscript{2}, is however barely detected; its concentration only reaches up to 0.98 at 0.91–1.04, see Table S 6. This syngas ratio is too low for optimal Fischer-Tropsch synthesis, where a ratio of 2 is desired [48]. Ethane, ethene, and propene show similar behaviour as seen with CH\textsubscript{4} reforming, i.e. a low–high–low concentration profile, but at lower values; only the ethene concentration surpasses the 1% mark, reaching 1.96% at 10 s and decreasing to 1.04%. Ethanol, the sole oxygenated hydrocarbon that could be quantified, is only formed in small amounts from 300 to 900 ppm.

Although some changes can be observed in ratios of different products, adding a SiO\textsubscript{2} packing to the reactor does not have a large impact on the product formation. In Fig. 5(b) we see that the H\textsubscript{2} concentration reaches the same end value, although it does so faster due to the higher overall rate coefficient, but a slightly larger amount of CO is formed at equilibrium (42.7% compared to 37.4%). As a result, the H\textsubscript{2}:CO ratio slightly changes to 0.98 at the shortest residence time, and to 0.94 at the longest. The other components remain almost constant.

Changing the CO\textsubscript{2}:CH\textsubscript{4} ratio has a large impact on the product formation, as can be seen in Fig. 5(c and d). A larger fraction of CO\textsubscript{2} in
the mixture (3:1 ratio) increases the CO fraction after a short residence time, but eventually reaches the same maximum concentration as for the 1:1 ratio, i.e. 37.1% vs. 37.4%, respectively. The H2 concentration, on the other hand, is significantly reduced over the entire residence time by almost a third, reaching a maximum of 23.8%, compared to 39% for the 1:1 ratio. The H2:CO ratio therefore decreases as well, to values between 0.49 and 0.64. The higher CO2 fraction leads to a slight increase in the O2 concentration, which is 0.008% at the shortest residence time and 0.0114% at the longest, but is still very low compared to pure CO2 splitting, as explained above. The concentration of the remaining components strongly decreases to values well below 1%, as can be seen in Table S6.

Vice versa, a large fraction of CH4 in the mixture (1:3 ratio) seems to be more beneficial overall. The CO concentrations are half the values reached for the 1:1 ratio across the entire range of residence times, but the H2 concentration increases by a factor 1.2 to 1.5, showing a maximum of 56.1%, and thus resulting in large H2:CO ratios between 2.34 and 3.08, which is more optimal for the production of (oxygenated) alkanes and alkenes via the Fischer-Tropsch process [48]. The ethane, ethene and propane production rises as well, with concentrations up to 4.02%, 0.32% and 1.43%, respectively, at 17.5 s. Higher CH4 fractions are however more susceptible to carbon depositions, requiring periodic physical cleaning, or chemical cleaning by e.g. pure O2 or H2. For this reason it could be interesting to add more CO2 in the mix, to prevent carbon deposition.

An interesting observation can be made for the CO and CO2 concentrations. For all DRM experiments—except at a 1:3 ratio and at long residence times, when most condensable products are being made—the sum of CO2out and CO2out after the experiment is equal to CO2in before the experiment, as can be seen in Table S6. This means that in DRM, on a global scale, all net CO2 conversion is dominated by the transformation of CO2 into CO, without further reaction towards hydrocarbons. It might be possible that on the molecular scale some CO reacts towards elemental C and O, or towards C1 oxygenates, and some hydrocarbons (or elemental C) partially oxidize to CO. However, on a reactor scale, CO2, besides mainly being a source of non-reactive CO, seems to act only as an oxygen source for the oxygenated hydrocarbons and water, which means that CH4 is expected to be the actual carbon source for the production of the (oxygenated) hydrocarbons. A comparative study to partial oxidation of CH4 (CH4 + O2 → C2H4O2) might be of interest to check the conversion efficiency (both on a rate coefficient and energy basis) towards (oxygenated) hydrocarbons.

Also, we observe that the produced fraction of CO in the empty reactor during CO2 dissociation matches the maximum produced fractions of CO in DRM with CO2 - CH4 fractions of 3:1 and 1:1. As the CO2 fraction in the mixture is reduced, the effective CO2 conversion increases, to maintain the same overall CO2 consumption, and thus CO production. It seems that the equilibrium concentration of CO2 in the empty reactor, in both CO2 dissociation and DRM, is limited to the same power-dependent maximum value of 36%, independent of the DRM ratio. Only when lowering the initial CO2 fraction to 25%, as in the 1:3 DRM ratio, we obtain lower fractions of CO, which is logical due to the low initial concentration of CO2. Hence, it confirms our conclusion of no noticeable production of CO from CH4. Also, when adding the SiO2...
packing material, we observe a higher CO production, due to the optimized kinetics, as discussed in section 3.3, although the maximum CO fraction here does not match the maximum CO fraction from SiO2-packed CO2 dissociation.

3.6. Further considerations

3.6.1. Strengths and limitations of this PCE study

The focus of this study is to accurately quantify the amount of (reacted) CO2 and CH4, as it is the input of the PCE model equation. Characterizing the other components in the reactor exhaust gas is a more difficult matter. First of all, it is impossible to know all the possible products that can be formed in the reactor beforehand, let alone quantify them all. This is because we cannot calibrate for them or collect them (in the case of solid or liquid depositions) in a reliable way; up to C7 and C4-OH were detected but only few calibrated, see table S 2 in the supplementary material.

Although we cannot determine all components, we can still correctly quantify all components for which the GC was calibrated because we measured no significant gas expansion nor contraction in the DRM and CH4 reforming experiments; in the case of pure CO2 dissociation we did measure gas expansion consistent with the reaction stoichiometry and this was corrected accordingly [25]. Therefore, the conversion of CH4 and CO2 could always be measured in a reliable way. As a result of the limited set of calibrated components, the atom balances in our experiments are not complete. Depending on the exact conditions, we missed up to 75% of a specific atom balance (most pronounced at low flow rates, SiO2 packing, and high methane content). This is for example visible in the pure CH4 experiments where the H2 concentration is on average 88% of that of the converted CH4 and thus most of the remaining CH2 from CH4 is missing and most likely will be deposited as liquid CxHy and solid C. A similar example can be seen for DRM: in section 3.5.3 we saw that, on a global reactor scale, CH4 acted like the sole carbon source for higher hydrocarbons as equimolar amounts of CO were formed compared to the reacted CO2. Still only a few extra (oxygenated) hydrocarbons and a minor amount of O2 could be quantified, resulting in incomplete C, H, and O balances. Table S 9 in the supplementary material shows all atom balances for all gas mixtures and for both empty and SiO2-packed reactors (if applicable).

Also, one can think about the validity of applying the partial chemical equilibrium (PCE) concept in these more complex chemical reactions (CH4 reforming and DRM), in comparison to CO2 dissociation. In the latter, CO2 is split into CO and O2, which are usually the only end products, unless significant O3 or carbon deposition would be present due to some conditions. Once the CO2 loss rate is equal to the formation rate by CO oxidation, PCE is reached and further extension of the residence time has no influence on the overall CO2 conversion, or CO and O2 formation. However, the chemistry in CH4 reforming and DRM is much more complex, which means that the manifestation of the PCE might be different for each component. In sections 3.1 to 3.4, we

Fig. 5. Measured concentration of different calibrated components for DRM, plotted as stacked bars as a function of residence time, for (a) the empty reactor with a CO2:CH4 ratio of 1:1, (b) the SiO2-packed reactor at 1:1 ratio, (c) the empty reactor at 3:1 ratio, and (d) the empty reactor at 1:3 ratio. The fitted DRM conversion is displayed as well, as a reference for each case. All components are measured on the TCD. The exact values of the concentrations can be found in Table S 6 in the supplementary material.
showed that the PCE appears to be reached in CH₄ reforming and DRM, based on the CO₂ or CH₄ conversions alone. An explicit proof for the existence of a global PCE state, however, would entail running the same conversion process starting from the pure products and verifying if this results in the same final gas composition, as we did in [25]. Such an approach is precluded here by the difficulty of identifying all species in these more complex gas mixtures. As noted earlier, smaller hydrocarbons might for example still polymerize to higher hydrocarbons, while the latter can dissociate back into lower hydrocarbons, and it is not clear from our results if true equilibrium was already reached in our experiments. For this reason, the PCE analysis in this work was mostly restricted to the easy-to-characterise initial reactants. At least for these gases, time-dependent concentrations appear to be consistent with PCE behaviour, in line with our previous more rigorous investigation of the chemically more simple CO₂ dissociation process [25]. As a result, we can confidently report PCE conversions of the feed gases, but not PCE yields for all the products. Additionally, despite the extensive nature of the (plasma) chemistry in CH₄ reforming and DRM, our results confidently prove that the overall gas conversion processes resulting from even these complex reactions can be described by our apparent first order kinetics PCE model, demonstrating opportunities towards other reaction and reactor types.

3.6.2. Interpreting the energy cost

Finally, we shortly reflect on the efficiency of the reactor configurations used in this work. For this, we invoke the concept of the energy cost (EC), i.e. the amount of energy necessary to convert one mole of reactant mixture, according to:

$$EC = \frac{SEI \cdot V_m}{\dot{V}_{\text{meas}}}$$  \hspace{1cm} (11)

with SEI the specific energy input based on the ratio of the plasma power and $\dot{V}$ the volumetric flow rate ($SEI = P/\dot{V}$), and $V_m$ the molar volume (22.4 L/mol). The calculated EC for all experiments can be found in Table S1 in the supplementary material.

The fact that the plasma conversion processes under consideration can be characterized by PCE behaviour has import consequences for their energy cost. The conversion asymptotically reaches the equilibrium conversion as an upper limit, which cannot be further improved by increasing the specific energy input (for a given set of plasma conditions). As a result, accurate determination of the location of the PCE, and the rate of evolution towards it, requires running conversion experiments at long residence time, resulting in very low energy efficiencies. However, such experiments would only have to be carried out once for a given process/reactor combination. Afterwards, the obtained information can be used to design or select an optimal process or conditions, determined by economical or practical considerations. More specifically, the rate coefficient and equilibrium conversion allow to predict the conversion as a function of residence time, which can be contrasted with energy cost at the same residence time; an optimal balance between the two can be selected, and compared with another process.

From the calculated EC data we can see that the minimum EC for almost all cases is reached at the shortest residence time (or highest flow rate), except for pure CH₄ reforming. This is due to the fact that, generally, the conversion rises too slowly with increasing residence time, or in other words, the rate at which the flow rate decreases cannot be matched by the increase in total conversion. For CO₂ dissociation we find a minimum EC of 1.7 kWh/mol at 2.9 s, while the minimum EC of CH₄ reforming is even higher, i.e. 2.6 kWh/mol at 10 s. In both cases, adding a packing material to the reactor does not yield a lower EC. Interestingly, DRM in a 1:1 ratio shows better results, with a minimum EC of 1.16 kWh/mol at 2.9 s. Both the 3:1 and 1:3 ratios perform slightly worse, with values of 1.32 and 1.38 kWh/mmol, respectively. From these data we conclude that the most effective use of the DBD plasma reactor is at shorter residence times. This conclusion might change however if we take other aspects into account, such as separation processes and the (liquid) higher hydrocarbons. Indeed, the separation of a low converted gas mixture might suffer from high running costs, rendering less energy efficient conditions but with higher overall conversion. An optimum can probably be found here, depending on the physical process volume and process parameters. Moreover, we found in sections 3.5.2 and 3.5.3 that the longer residence times produced higher amounts of products, and also more diverse products. Depending on the desired end products, i.e. either syngas, lower hydrocarbons, or liquids, different operating conditions might be preferred. Therefore, a careful analysis of this type of experiments is very useful for the design of optimised processes for specific purposes.

4. Conclusion

In this paper we investigated the kinetics of the CO₂ dissociation and CH₄ reforming reactions and how they influence each other when both gases are combined in DRM. Fitting the time evolution of the gas composition to a new generalised first order kinetic model for the partial chemical equilibrium made it possible to determine multiple trends, elucidate macroscopic changes in the plasma chemistry, and link them to changes in the loss or formation reactions of the reagents. CO₂ dissociation exhibits a higher apparent rate coefficient (0.120 s⁻¹) than CH₄ reforming (0.041 s⁻¹), but CH₄ reforming has a higher equilibrium conversion (82%) than CO₂ dissociation (53.6%). The lower rate coefficient of CH₄ reforming is attributed to fast dissociation and recombination reactions, rendering a slow net overall rate. Mixing both gases in a 1:1 ratio combines the best of both worlds, i.e. the higher equilibrium conversion of CH₄ reforming, yielding 75.4%, and the higher rate coefficient of CO₂ dissociation, ending up at 0.088 s⁻¹. These results point to additional interactions of the two gases, which open new pathways by the individual gas products.

Adding a spherical non-porous SiO₂ packing material to the reactor increases the electron temperature and thus further stimulates electron impact-based processes, causing gas-specific effects. In a pure CO₂ plasma, the equilibrium shifts further away to 71.1%, at the cost of slightly reducing the overall conversion rate to 0.111 s⁻¹, whereas for CH₄ reforming the equilibrium conversion stays about the same at 81% and the apparent rate coefficient increases to 0.074 s⁻¹. Mixing the gases results in an increase of both equilibrium and rate coefficient in 1:1 DRM to 84.3% and 0.130 s⁻¹, respectively.

Comparing different CO₂:CH₄ ratios reveals the delicate balance of the combined chemistry. CO₂ drives the loss reactions in DRM, resulting in higher reaction rate coefficients when present in higher fractions; the presence of CH₄ in the mixture suppresses back reactions, resulting in higher equilibrium conversions when it is more abundant.

Finally, analysis of the effluent of all experiments revealed not only how the product composition changes and is influenced by mixing the CO₂ and CH₄, but also how it changes in time. We see trade-offs between producing larger amounts of hydrocarbons (when more CH₄ is present in the mixture), and optimal H₂:CO syngas ratios (when more CH₄ is present).

The method for kinetic analysis used in this work is shown to be a practical way to describe the plasma-based conversion of molecules of interest in arbitrary gas mixtures with a priori unknown stoichiometry. In addition, we have shown how analysis of the derived kinetic parameters can elucidate key mechanistic aspects of the conversion process and help bridge the gap with detailed kinetic models. We therefore highly advise this method for any further research in plasma (catalysis) based gas conversion.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to


References


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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.cej.2020.126630.