Gliding arc plasma for CO\textsubscript{2} conversion: Better insights by a combined experimental and modelling approach

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ABSTRACT

A gliding arc plasma is a potential way to convert CO\textsubscript{2} into CO and O\textsubscript{2} due to its non-equilibrium character, but little is known about the underlying mechanisms. In this paper, a self-consistent two-dimensional (2D) gliding arc model is developed, with a detailed non-equilibrium CO\textsubscript{2} plasma chemistry, and validated with experiments. Our calculated values of the electron number density in the plasma, the CO\textsubscript{2} conversion and energy efficiency show reasonable agreement with the experiments, indicating that the model can provide a realistic picture of the plasma chemistry. Comparison of the results with classical thermal conversion, as well as other plasma-based technologies for CO\textsubscript{2} conversion reported in literature, demonstrates the non-equilibrium character of the gliding arc, and indicates that the gliding arc is a promising plasma reactor for CO\textsubscript{2} conversion. However, some process modifications should be exploited to further improve its performance. As the model provides a realistic picture of the plasma behaviour, we use it first to investigate the plasma characteristics in a whole gliding arc cycle, which is necessary to understand the underlying mechanisms. Subsequently, we perform a chemical kinetics analysis, to investigate the different pathways for CO\textsubscript{2} loss and formation. Based on the revealed discharge properties and the underlying CO\textsubscript{2} plasma chemistry, the model allows us to propose solutions on how to further improve the CO\textsubscript{2} conversion and energy efficiency by a gliding arc plasma.

1. Introduction

Plasma technology offers unique perspectives, because of its capacity to induce chemical reactions within gases at ambient temperature and pressure, due to its non-equilibrium character. Plasma is created by applying electric power to a gas, causing breakdown of the gas into ions and electrons and also producing a large number of reactive species, such as various radicals and excited species. This makes plasma a highly reactive cocktail, which is quite promising for greenhouse gas conversion. Indeed, the inert CO\textsubscript{2} gas is activated by electron impact ionization, excitation and dissociation. Furthermore, plasma is very flexible and can easily be switched on and off, so it is quite promising for storing peak renewable energy into fuels. Indeed, more and more electrical energy nowadays is produced from renewable energy sources (wind or solar), which often suffer from fluctuating peak powers, making it difficult to match the supply of this electricity with the demand. This surplus of electricity can in principle be used in plasma to convert greenhouse gases into value-added chemicals when adding a suitable H-source to the CO\textsubscript{2} gas, such as H\textsubscript{2}O, CH\textsubscript{4} or H\textsubscript{2}. However, there is still a long way to go, certainly if we target the selective production of some value-added products, for which the combination with a suitable catalyst would be needed. This makes plasma based greenhouse gas conversion fit in principle in the framework of green chemistry [1,2] and also complies with the “cradle-to-cradle” principle [3].

Gliding arc (GA) plasmas are potential plasma sources for gas conversion [4–18] because they offer benefits of both thermal and non-thermal discharges. They are typically considered as ‘warm’ discharges, which are characterized by a better energy efficiency than other types of plasmas. The reason is that they provide efficient vibrational excitation of the molecules, which is seen as the most energy-efficient way to split CO\textsubscript{2} molecules [19].

In order to improve the applications (i.e., mainly gas conversion), the physical and chemical characteristics of the GA have been extensively studied by experiments, including high-speed photography [20], electrical measurements [21–23] and spectroscopic measurements [24,25]. Besides experiments, detailed modelling is also very useful to provide more insight into the underlying reaction mechanisms of plasma assisted gas conversion or synthesis, not only in a GA but also in other types of plasmas. For example, computer modelling is widely used to evaluate quantities which are difficult to measure, and to
identify the most important chemical reactions [26–32]. However, only a few papers in literature deal with modelling of a GA, typically applying a 1D analytical model, such as the Elenbaas–Heller model [33] or the plasma string model [34] without a detailed description of the chemical reactions occurring in the GA. Recently, a 2D non-quasi-neutral model was presented to study the arc root movement in an argon GA [35,36]. Moreover, 3D quasi-neutral models for a novel type of GA plasmatron [37] and a classical diverging electrode GA reactor [38,39] were also reported. However, these models were all developed for argon. For a GA operating in CO2, the large number of species and related chemical reactions makes spatially resolved models computationally expensive. That is why only a limited number of numerical studies were reported so far on this subject, with only two papers for GA based CO2 conversion published to

It is clear that more research is needed to fully exploit the capabilities of the GA for CO2 conversion. In this paper, we therefore present a combined modelling and experimental study, based – for the first time – on a 2D model. The aim of this study is not only to elucidate the underlying mechanisms, but also – based on the obtained insights – to propose solutions on how to further improve the performance of the GA for CO2 conversion.

2. Experimental setup of the GA reactor

Fig. 1 illustrates the experimental setup of the GA and surrounding measurement equipment. The GA reactor consists of two stainless steel semi-ellipsoidal electrodes with thickness of 2 mm (60 mm long and 18 mm wide) fixed in an insulating bracket and symmetrically placed on both sides of a gas nozzle with a diameter of 1.5 mm. The reactor is designed to facilitate easy electrode replacement, and the discharge gap between both electrodes, as well as the distance between the nozzle exit and electrode throat, is adjustable. Pure CO2 gas was injected into the GA reactor and it pushes the arc plasma, which is initiated at the shortest gap between both electrodes, towards larger interelectrode distance until it extinguishes, and a new arc is created at the shortest discharge gap between the two electrodes as well. 2 mm. The arc dynamics are revealed by means of a digital high-speed camera (Phantom V.7.1) which can record up to 4800 pictures per second using the full 800 × 600 pixel SR-CMOS imaging sensor array. The measurement technique was intensely optimized to fine-tune the best recording conditions. The frame rates to visualize the arc propagation and the exposure time of the detector to enhance the contrast between the arc and the reactor were investigated. The products of the CO2 conversion after passing through the GA reactor were sampled when the plasma reaction has reached a stable condition, i.e., typically after 30 min. The gaseous products were analyzed by a gas chromatograph (Shimadzu, GC-2014) equipped with a thermal conductivity detector (TCD) and a flame ionization detector (FID). As we mention below, a standard case of 2.5 L/min and 40 W is used to validate our model. Furthermore, the vertical distance between the nozzle exit and electrode throat was 2 mm and the shortest discharge gap between the two electrodes was also 2 mm.

The plasma power is calculated by integration of the arc voltage and current, as shown in Eq. (1).

\[ P_{\text{plasma}} = \frac{1}{T} \int_0^T V_{\text{plasma}} \times I_{\text{plasma}} \, dt \]  

(1)

The conversion of CO2, \( X_{\text{CO2}} \), is defined as:

\[ X_{\text{CO2}}(\%) = \frac{CO_{2(\text{in})} - CO_{2(\text{out})}}{CO_{2(\text{in})}} \times 100\% \]  

(2)

where \( CO_{2(\text{in})} \) and \( CO_{2(\text{out})} \) are the CO2 signals without and with plasma, respectively. Since the method mentioned above does not account for the gas expansion due to CO2 splitting, a correction factor is used, which is explained in the supplementary information of Ref. [31].

In order to calculate the energy efficiency of CO2 conversion, the specific energy input (SEI) in the plasma is defined as:

\[ SEI\left(\frac{kW}{L}\right) = \frac{\text{Plasmapower (kW)}}{\text{Flowrate}(\frac{L}{min}) \times 60}\left(\frac{s}{min}\right) \]  

(3)

where the flow rate is expressed in L/min (liters normal per minute) with reference conditions at a temperature of 0 °C and a pressure of 1 atm.

The energy efficiency, \( \eta \), is calculated as:

\[ \eta(\%) = \frac{\Delta H_f \left(\frac{\text{kJ}}{\text{mol}}\right) \times X_{\text{CO2}}(\%)}{SEI\left(\frac{\text{kJ}}{\text{L}}\right) \times 22.4 \left(\frac{\text{L}}{\text{mol}}\right)} \]  

(4)

where \( \Delta H_f \) is the reaction enthalpy of CO2 splitting (i.e., 279.8 kJ/mol), \( X_{\text{CO2}} \) is the amount of CO2 converted, SEI is defined above and 22.4 L/mol is the molar volume at 0 °C and 1 atm.

The experiments were performed 4 times and they were reproducible within ± 5% of the averaged values.

Fig. 1. Schematic diagram of the GA experimental set-up.
3. Description of the 2D plasma slab model

3.1. The GA reactor geometry

The 2D fluid model that we developed applies to a Cartesian geometry, which allows to describe the gliding of a ‘2D arc’, which is basically a finite plasma slab. The simulated geometry in the direction perpendicular to the simulation plane is assumed to be equal to the electrode thickness of 2 mm. Hence, the electrical current in the 2D model is obtained by integration of the current density over the arc slab, which fits the experimental signal. Furthermore, the flow field is determined by taking into account a flow passing channel with a depth of 2 mm with the specified flow rate. In this way, the calculated gas velocity is similar to the experimental data when the vertical distance between the nozzle exit and electrode throat was 2 mm and the shortest discharge gap between the two electrodes was also 2 mm. Indeed, a rough estimation of the experimental gas velocity is obtained by examination of the arc displacement shown in successive high-speed photographs (see supporting information). In principle, a 3D model would be required to describe the GA behaviour in a realistic way, in view of the intrinsic 3D nature of the GA. However, a 3D model is very time consuming and it requires significant computer resources, especially when modelling a CO2 plasma with complicated plasma chemistry. Furthermore, previous work for an argon GA [38] has shown that the results of a 2D model compare well with those of a 3D model, and can thus be used for a better understanding of the GA basic characteristics. The total width and height of the model geometry, including the region outside the electrodes where the gas can flow without passing through the arc, is 38 mm and 70 mm, respectively.

3.2. CO2 plasma chemistry and treatment of the vibrational levels

The chemistry set is based on the full chemistry set developed by Kozák and Bogaerts [26,27] with a 0D model, but reduced to include only the most important species and processes. In this way, we can avoid excessive calculation times in this 2D model, but we still account for the vibrational kinetics, which is crucial for describing CO2 conversion in a GA plasma reactor [41]. The list of species considered in the model is shown in Table 1. These species include various neutral molecules in the ground state, as well as in various electronically and vibrationally excited levels, a number of radicals, positive and negative ions, and the electrons. In the full model of Kozák and Bogaerts [26,27], 25 CO2 vibrational levels (i.e., 4 effective levels of the symmetric modes and 21 levels of the asymmetric stretch mode, up to the dissociation limit) were taken into account. However, to further reduce the calculation time, which is needed to implement this chemistry in a 2D model, Berthelot and Bogaerts [42] developed a level lumping method, which groups the 21 asymmetric stretch mode vibrational levels into a number of lumped levels, without loss of essential information. We applied this level lumping method in [41] for a 1D gliding arc model, and we illustrated that lumping the 21 levels into 3 groups can reproduce the plasma properties, the vibrational distribution function (VDF) and the CO2 conversion very well. Therefore, we adopt here the same level lumping method with 3 groups for the asymmetric stretch mode, with each group including 7 vibrational levels (group 1: CO2[v1-v7], group 2: CO2[v8-v14], group 3: CO2[v15-v21]). The species number density of each level within one group can be determined following the method described in [41,42]. Besides, we also take into account the 4 effective levels of the symmetric modes (CO2[v1] - CO2[v4]), 1 electronically excited level of CO2 (CO2[e]), and 3 vibrational levels of O2 (O2[v1] - O2[v3]), as indicated in Table 1.

All these species undergo a large number of chemical reactions, i.e., electron impact collisions with neutral species, leading to excitation, ionization, dissociation and electron attachment, electron-ion recombination reactions, as well as many heavy-particle chemical reactions (i.e., ion-ion, ion-neutral and neutral-neutral reactions). We pay special attention to the reactions of the vibrational levels, i.e., electron impact vibrational excitation, and vibrational energy exchange upon collision with ground state species or other vibrationally excited levels (i.e., so-called vibrational-translational (VT) and vibrational-vibrational (VV) relaxation, respectively). Moreover, the same chemical reactions as for the ground state species are carefully included for the vibrational levels as well, because the vibrational energy can help overcome the activation energy barrier of the reactions and thus increase the reaction rate of CO2 splitting. The chemical reactions, the corresponding rate coefficients and the references where these data were adopted from, are listed in our previous work [41].

3.3. System of governing equations and boundary conditions

The model calculates the densities of all the plasma species, the electron temperature and gas temperature and the electric field in the GA, as well as the gas flow profile. We assume electrical neutrality in the plasma, because the sheath is not considered in our model. This assumption has no significant influence on the arc column [39]. The species densities and the electron mean energy are calculated with continuity equations based on transport and on production and loss terms defined by the chemical reactions (and by Joule heating for the electron energy). The species transport is based on drift in the electric field and diffusion due to concentration gradients. As we assume electrical neutrality in the arc plasma, the ambipolar electric field is calculated from the charged species densities. The gas heat transfer equation is solved for the gas translational temperature, and finally, the neutral gas flow, which is responsible for the arc displacement, is described by the Navier-Stokes equations, providing a solution for the mass density and the mass-averaged velocity. The Navier-Stokes equations are first solved separately, and subsequently, the obtained velocity distribution is used as input data in the other equations, describing the plasma behaviour and the gas heating. The equations solved, as well as the corresponding boundary conditions, are explained in detail in the supporting information. Finally, the external circuit and the power supply need to be specified in the simulation. The source voltage has a sinus shape, \( V_{\text{source}} = 7200 \sin(2\pi 50 + 0.50) \text{ V} \), and a resistance of 60 k\( \Omega \) is used to limit the discharge current; it provides a total arc discharge power of 40 W, which is similar to the typical experimental value at a gas flow rate of 2.5 L/min.

The equations are solved by means of the COMSOL Multiphysics software [43], a commercial finite element software designed for solving problems of multi-physics. As initial values we assume that the concentrations of CO2 in the ground state and in the various excited levels follow a Maxwellian distribution at room temperature.

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Overview of the plasma species included in the model.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neutral ground state species</td>
<td>CO2, CO, C, O2, O</td>
</tr>
<tr>
<td>Neutral excited states</td>
<td>CO2[v1], CO2[v2], CO2[v3], CO2[v4], CO2[v5], CO2[v6], CO2[v7], CO2[v8], CO2[v9], CO2[v10], O2[v1], O2[v2], O2[v3]</td>
</tr>
<tr>
<td>Charged species</td>
<td>CO2^+, O2^+, CO2^−, O^+, O2^−, e^-</td>
</tr>
</tbody>
</table>

4. Results and discussion

In section 4.1 we will first validate our model by comparing our calculated values with experimental data for the electron number density (which is one of the most important plasma properties), as well as for the CO2 conversion and corresponding energy efficiency. Subsequently, in section 4.2 we will benchmark our results for the CO2 conversion and energy efficiency to the classical thermal conversion process and to other plasma-based technologies for CO2 conversion.
reported in literature. This allows us to provide a clear overview of the capabilities of the GA for CO₂ conversion, as well as its limitations, for which we should propose some process modifications, to further improve the results. In order to achieve this, we need a better insight in the typical discharge characteristics, as calculated by the model, which will be presented in section 4.3. Furthermore, we will also perform a chemical kinetics analysis in section 4.4, to elucidate the role of various plasma species and their reactions in the GA based CO₂ conversion. Finally, based on the revealed discharge properties and the obtained plasma chemistry, we will propose in section 4.5 some solutions on how to further improve the CO₂ conversion and the energy efficiency by the GA.

### 4.1. Experimental validation of the model

In Table 2 we compare our calculated results for the electron number density, CO₂ conversion and corresponding energy efficiency, with the corresponding measured values, at a typical experimental gas flow rate of 2.5 L/min and a discharge power of 40 W.

<table>
<thead>
<tr>
<th>Results</th>
<th>Electron number density</th>
<th>Conversion</th>
<th>Energy efficiency</th>
</tr>
</thead>
<tbody>
<tr>
<td>Calculation</td>
<td>$10^{18}$–$10^{19}$ m⁻³</td>
<td>2.78%</td>
<td>32.8%</td>
</tr>
<tr>
<td>Experiment</td>
<td>$2.6 \times 10^{19}$ m⁻³</td>
<td>2.90%</td>
<td>34.3%</td>
</tr>
<tr>
<td>Experimental error</td>
<td>4.9%</td>
<td>4.3%</td>
<td>4.6%</td>
</tr>
</tbody>
</table>

\[
\langle n_e \rangle = \frac{\sigma}{\epsilon_e} \tag{6}
\]

With \(\epsilon\) the electron charge. Using a time averaged gas temperature of 2400 K and an electron temperature of 1.7 eV, as obtained from our model (see section 4.3), we calculated \(\mu_e = 0.56\) m²/V·s by means of a Boltzmann equation solver BOLSIG+ [44]. Hence, formula (6) gives an estimate of the time and spatially averaged electron number density, \(\langle n_e \rangle \approx 2.6 \times 10^{18}\) m⁻³. Our calculations predict the maximum electron number density in the discharge channel to be around $10^{19}$ m⁻³ (see section 4.3). Considering the non-uniform distribution within the discharge channel, we can obtain a spatially averaged value of the electron number density within the range $10^{18}$ m⁻³–$10^{19}$ m⁻³, indicating a reasonable agreement between the calculated and measured values.

The calculated conversion of CO₂, \(X_{\text{CO}_2}\), is determined as:

\[
X_{\text{CO}_2} = \frac{\int_{\text{CO}_2} Q \, dt}{\int_{\text{CO}_2} \, dt} \times 100\% = \frac{Q_{\text{CO}_2}}{Q_{\text{CO}_2}} \times 100\%
\]

where \(Q_{\text{CO}_2}\) is the particle flow rate of CO₂ entering the reactor per second (in s⁻¹), \(f_{\text{CO}_2}\) is the net splitting rate of CO₂ inside the arc (in m⁻³ s⁻¹), and \(l_0 = 2\) mm, is the thickness of the GA reactor (see below).

The particle flow rate of CO₂, \(Q_{\text{CO}_2}\), represents the total number of CO₂ molecules flowing into the reactor per second, and is obtained as follows:

\[
Q_{\text{CO}_2} = \frac{k \, (\frac{2}{k})^\frac{3}{2}}{P_0} \times \frac{\mu_0}{\epsilon_0} \times \frac{1}{\epsilon} \times \frac{1}{\mu} \times \frac{1}{\hbar} \times \frac{P_0}{k_0(T_0)} \times 100\%
\]

where \(k\) is the Boltzmann constant, \(Q_a\) is the gas flow rate at the standard temperature \(T_0 = 273\) K and pressure \(P_0 = 101,325\) Pa.

The net splitting rate of CO₂, \(r_{\text{CO}_2}\), in m⁻³ s⁻¹, represents the net number of dissociated CO₂ molecules per volume and per second, and is obtained by taking into account all the chemical reactions, leading to destruction (when a positive value) or formation (when negative) of CO₂ molecules. In order to determine the total conversion of CO₂, as shown in Eq. (6), the net splitting rate of CO₂, \(r_{\text{CO}_2}\), is integrated spatially over the whole reactor and temporally over the whole gliding cycle. Because of the prohibitively long computation time in a 3D model, a 2D plasma slab model is used, assuming that the distribution of plasma parameters in the direction perpendicular to the simulation plane (see Fig. 2b) is uniform. As a result, the arc is not a "wire" but a "slab" with a length \(l_0\) in the direction perpendicular to the simulation plane.
plane. We assume \( l_0 \) is equal to the thickness of the GA reactor, i.e., 2 mm. Thus the total conversion of CO\(_2\) in the 2D model is obtained by the integration of the net splitting rate of CO\(_2\)-CO\(_2\) over the arc slab with \( l_0 = 2 \) mm.

Our calculated conversion and energy efficiency of CO\(_2\), at a gas flow rate of 2.5 L/min and a discharge power of 40 W, are 2.78% and 32.8%, respectively, which is also in satisfactory agreement with the experimental values of 2.90% and 34.3%. The comparison of these three key parameters indicates that our model most probably can provide a realistic picture of the plasma chemistry.

Comparison of other plasma characteristics, such as the electron temperature or gas temperature, was not possible, as the latter properties could not be determined in our experimental setup, and are also not available in literature for a pure CO\(_2\) GA. This is probably because optical emission spectrometry is not suitable here, as there are no proper spectral lines that can be used. However, our calculated values for electron temperature (up to 1.7 eV) and gas temperature (up to around 2700 K) are comparable with experimental data from literature, for GA reactors using other molecular gases (nitrogen and air) [45–47], as well as for gaseous mixtures containing CO\(_2\) [48,49]. For example, Wu et al. [48] measured values for the electron excitation temperature of approximately 1.1–1.7 eV, using a rotating GA reactor for a mixture of CH\(_4\)/CO\(_2\). Moreover, in a non-equilibrium GA “tornado” discharge using CO\(_2\) doped with 1% N\(_2\), the rotational gas temperature was determined to be 2700 K \( \pm 50 \) K [49].

We can only compare here the calculated and experimental data at a gas flow rate of 2.5 L/min and discharge power of 40 W, because at these conditions the arc was observed to glide smoothly along the electrodes. Indeed, at higher gas flow rates, a phenomenon of back-breakdown occurs, affecting the arc gliding process (see further). These back-breakdown events cannot self-consistently be captured by the model, because this behaviour is mostly stochastic by nature and the arc instabilities are not well defined. Therefore, we would need to make some assumptions in the model on the number of back-breakdown events, and depending on the values assumed for the back-breakdown frequency, we would always be able to obtain good agreement with the experiments. Hence, we lose the real validation possibility at higher gas flow rates. Therefore, we could only validate the model at a gas flow rate of 2.5 L/min and a discharge power of 40 W, where our high speed camera did not record any back-breakdown events. However, in section 4.5, we will assess the effect of a different number of back-breakdown events on the calculated conversion and energy efficiency, which can in principle be correlated with different values of gas flow rate and discharge power.

4.2. Comparison of our results with other plasma systems from literature

In Fig. 3, we compare our results for the energy efficiency vs CO\(_2\) conversion with data obtained from literature for CO\(_2\) splitting, in other GA discharges [5,6,50,51], as well as in other types of plasma reactors, such as microwave (MW) plasma [52–55], dielectric barrier discharge (DBD) [28,56–60], nano-second pulsed plasma (NSPP) [61,62], corona discharge [63,64], micro hollow cathode discharge (MHC) [65,66] and spark discharge [67]. We can conclude that in terms of energy efficiency, the GA plasma is very promising, similar to the corona discharge [63,64]. It should be mentioned that for MW plasmas some higher energy efficiencies (i.e., up to 80 and 90%) were obtained in literature by Rusanov et al. [68] and Asisov et al. [69], respectively. However, their MW plasma reactors were operating at a reduced pressure of 0.06–0.26 atm and 0.05 – 0.2 atm, respectively, and thus they need vacuum equipment. This makes it more difficult to be applied on industrial scale, and the energy cost of the pumping system should also be included when calculating the energy consumption. Bongers et al. recently obtained values up to 50%, when applying a reverse vortex gas flow [70], but again these experiments were conducted at reduced pressures of 150–600 mbar (0.15–0.60 atm). In order to allow a fair comparison, we therefore only present results in Fig. 3, obtained at atmospheric pressure. When the MW discharge is operating at atmospheric pressure, the reported energy efficiency dramatically drops to values of about 5–20% [52–55].

If we compare our results with those obtained in other GA reactors from literature, it is important to explain that there exist roughly two different reactor designs. The classical GA reactor, which is used in this study, typically consists of two plane diverging electrodes between which the gas flows. In contrast, recently a three-dimensional GA reactor, consisting of cylindrical electrodes with tangential gas inlet, leading to a vortex gas flow configuration, has been developed, also called GA plasmatron (GAP) [19]. Indarto et al. [5] applied a classical GA configuration, like in our case, and they obtained a highest energy efficiency of around 17%, which is much lower than our current work. On the other hand, Nunnally et al. [6], Liu et al. [50] and Ramakers et al. [51] used a vortex flow GAP, which can reach a somewhat higher conversion and energy efficiency. This reactor design is indeed very promising, because it can be more easily implemented in industry and the specific gas flow configuration ensures the gas treatment to be more uniform. This indicates that a better design of the classical GA reactor, to enhance the treated gas volume, would improve the conversion performance, as will be discussed in detail in section 4.5 below. However, in general we can deduce from Fig. 3 that the GA plasma shows a very good performance with a relatively high energy efficiency. This is because the energy efficient vibrational excitation processes are favoured, as will be revealed in section 4.4 below.

It is obvious from Fig. 3 that a DBD plasma [28,56–60] has a reasonable conversion but a quite low energy efficiency. This is due to the non-ideal operating conditions, as the electron temperature is typically higher than in a GA (or MW) plasma [19,71], and the mechanism of CO\(_2\) conversion involves charged and electronically excited species, and thus it is limited by the high energy cost for the formation of these species. The same applies for the nano-second pulsed plasma (NSPP) [61,62] which also has a rather low energy efficiency. The process capability of the micro hollow cathode discharge (MHC) [65,66] is very limited due to its very small volume. Therefore, it generally also exhibits a relatively low energy efficiency. The spark discharge [67] has a very high conversion, because of the very high energy consumption. The energy efficiency is also quite high, but it is lower than the thermal conversion process. This may be attributed to the fact that most of the energy is spent on the gas heating and the energy exchange with the surroundings. In general, we can conclude that the energy efficiency in our GA reactor at atmospheric pressure is better than the DBD plasma, microwave plasma, nano-second pulsed plasma and micro hollow cathode discharge plasma, and comparable to the corona discharge [63,64].

Finally, we also benchmark our results for the GA based CO\(_2\) conversion to the pure thermal conversion process (see the calculation method for the latter in the supporting information). It is clear that the CO\(_2\) conversion in our GA proceeds more energy efficient than pure thermal conversion. This is because the energy in the thermal conversion is distributed over all degrees of freedom based on the equipartition principle of energy, and thus it is especially spent on gas heating rather than on CO\(_2\) dissociation reactions. In contrast, our GA clearly operates in non-equilibrium conditions, as the electrons have a much higher temperature than the gas itself (see our calculation results in section 4.3 below). These highly energetic electrons induce different chemical reactions, which normally do not occur at the considered gas temperature in case of equilibrium conditions.

In spite of the reasonable results obtained already by the gliding arc, the conversion should still be further improved, while maintaining the high energy efficiency. More specifically, if this low conversion could not be further improved, it would imply the need for operating in a recycle mode, which would make the system highly non-effective.
4.3. Typical GA discharge characteristics

In order to understand the time behaviour of the plasma characteristics in the CO₂ GA, we plot in Fig. 4 the electron number density, electron temperature, gas temperature, as well as of the CO molar fraction distribution, at different moments in time, for a gas flow rate of 2.5 L/min and a discharge power of 40 W.

The results are plotted starting from $t = 1 \text{ ms}$. At $t = 0 \text{ ms}$, the source voltage is larger than the critical breakdown voltage with a shortest gap separation of 2 mm. The discharge ignition takes place, because of a positive value of the net electron generation, yielding an abrupt increase of the electron number density during the electrical breakdown. Once the conducting channel is established, the arc travels along the electrodes as a result of the gas flow drag. Since the gas velocity has a maximum value at the discharge axis and gradually decreases to zero at the electrode surface, the arc root moves at a much slower velocity compared to the arc body. Thus, the arc gradually begins to bend due to the gas blast. The maximum electron number density also increases due to the rising voltage and hence discharge current (see Fig. S1 in the supporting information), till a peak value is reached at 3.5 ms (see Fig. 4(a)). At later times, the discharge current drop, and consequently, the electron density follows the same trend till zero at $t = 8.5 \text{ ms}$, when the applied voltage reaches zero (see Fig. S1). The GA gradually extinguishes and enters a relaxation stage, where the voltage is small and not enough to sustain the GA discharge. Thus, there is a decaying residual low density plasma moving downstream with the gas flow (see Fig. 4(a)). Shortly after $t = 8.5 \text{ ms}$, the applied voltage of the alternating current (AC) power source changes its polarity (see Fig. S1 of the supporting information) and reaches again the critical breakdown voltage at the narrowest electrode gap separation of 2 mm, where a restrick occurs by establishing a new conducting channel. It should be noted that the re-ignition of the GA does not exactly take place at the shortest gap separation ($Y = 2.5 \text{ mm}$), but at $Y = 7.5 \text{ mm}$. This is because the local electric field at $Y = 7.5 \text{ mm}$ first reaches the critical breakdown field. This is in good agreement with our experiments, recorded by the digital camera (see Fig. S2 of the supporting information).

The rise and drop in electron number density during one GA discharge cycle results in an enhanced and reduced Joule heating effect before and after $t = 3.5 \text{ ms}$, respectively. The Joule heating refers to the process by which the passage of an electric current through a conductive medium produces heat and causes heating of the electrons. Correspondingly, the electron temperature first increases and then decreases (see Fig. 4(b)). After $t = 8.5 \text{ ms}$, the electron temperature of the residual GA channel continuously decreases, because the electron number density and the electric energy stored in the channel decay very rapidly. Subsequently, the extremely large reverse polarity voltage imposed across the electrodes at the shortest electrode gap leads again to an increase of the electron temperature and hence a subsequent breakdown at the new position of $Y = 7.5 \text{ mm}$ (see Fig. 4(b)).

Once the discharge is ignited, the electrons cause vibrational excitation of CO₂, and the energy stored in the vibrationally excited states will partially be transferred to the gas by vibrational-translational (V-T) relaxation. Indeed, at atmospheric pressure, the typical characteristic time for V-T relaxation in CO₂ is very short (around $10^{-5} \text{ s}$). As a result, the gas temperature also rises as a function of time, reaching a maximum value of about $2700 \text{ K}$ at around $t = 3.5 \text{ ms}$, when the applied source voltage ($V_{\text{source}} = 7200 \sin(2 \pi 50 t + 0.50)$) and the discharge current reach their maximum (see Fig. 4(c)). Subsequently, the gas temperature in the arc channel gradually decreases to around $2000 \text{ K}$ when a new cycle starts at $t = 10 \text{ ms}$, because the discharge power decays rapidly in the relaxation stage from 8.5 ms to 10 ms. The CO molar fraction is obviously equal to zero before the arc is formed, but it starts increasing gradually as a function of time, when the voltage and hence the discharge current in the arc rise, up to a value of 0.55 at $t = 3.5 \text{ ms}$, indicating that CO₂ is gradually converted into CO. At later times, the discharge current and hence the discharge power start to drop, so the CO molar fraction within the arc channel gradually decreases until the arc is extinguished. This is caused by recombination of CO and O into CO₂. Furthermore, new CO₂ gas will continuously be transported into the arc channel by both diffusion and convection, while the dissociation products will leave the discharge channel by the same transport mechanisms. This leads to a reduction of the maximum local CO molar fraction, as is clearly indicated in Fig. 4(d). Note that the overall CO₂ conversion is much lower than the local conversion of 80%, which corresponds to the maximum CO molar fraction of 0.55 (and CO₂ molar fraction of 0.2; see below). This is because the overall CO₂ conversion is calculated for the entire gas passing through the reactor, integrated over the time of one GA cycle (i.e., 10 ms), and thus not only for the fraction of gas passing through the active arc channel at a certain moment in time.

The molar fractions of the major neutral and charged species occurring in the CO₂ GA are plotted as a function of Y position in Fig. 5, at a time instant of 2.5 ms, and at the same conditions as in Fig. 4. It is clear that CO₂ is the major component in the plasma, except at the centre of the arc, where the molar fraction of CO₂ (around 0.2) is lower than the fraction of CO (around 0.5), and comparable to the molar fractions of O₂ (0.16) and O (0.14). This indicates that the majority of CO₂ is split here into CO and O₂ as well as O atoms. Moreover, part of the O atoms have recombined into O₂ molecules, indicating a higher
decay rate of the O molar fraction than that of O₂. The molar fractions of CO, O and O₂ drop quickly when moving towards the outer part of the arc, indicating that most of the CO₂ splitting takes place in the centre of the arc.

The CO₂ conversion can be further enhanced when applying a higher power, however, even at 100 W, the local molar fraction of CO₂ drops to extremely low values and the local conversion in the GA reaches almost 100%. This limits the further improvement of GA based CO₂ conversion. Therefore, the conversion can only be further enhanced if we can provide more CO₂ into the arc centre, while at the same time remove the dissociation products (CO and O₂) out of the arc centre. This will be further discussed in detail in section 4.5.

The molar fractions of the various charged species are at maximum $10^{-5}$, even in the arc centre, and they clearly drop upon larger distance from the centre of the arc. Also the electron molar fraction is at maximum $10^{-5}$, indicating that the CO₂ plasma is only weakly ionized, even in the centre of the arc. The major positive ions are the $\text{O}_2^+$ ions, while the $\text{CO}_3^-$ ions are the major negative ions, and they are even more important (although still with very low molar fractions) than the electrons, except in the centre of the arc. These trends are in agreement with our previous findings obtained by a 1D cylindrical discharge model, despite the considerable number of approximations adopted there [41].

The gas temperature and electron temperature are also plotted in Fig. 5. They both reach their maximum in the centre of the arc, as is logical, and they drop significantly as a function of position from the arc.
The electron temperature reaches a maximum of 1.5 eV (or 17,400 K) in the centre of the arc at the time instant of 2.5 ms, but it drops significantly as a function of rising distance from the arc centre in the first 0.5 mm, followed by a slower decay to thermal values at a distance of about 1.0 mm from the centre. The gas temperature is at maximum about 2700 K in the centre of the arc. From the comparison between these temperatures, it is clear that the gliding arc is far from thermal equilibrium, as the electron temperature is about 6 times higher than the gas temperature. As mentioned in section 4.1 above, a gas temperature up to around 2700 K and an electron temperature up to 1.5 eV correspond well to experimental data found in literature for low current atmospheric pressure GA discharges, although it should be mentioned that it is not easy to compare different GA setups with different reactor geometries and discharge conditions.

4.4. CO2 conversion mechanisms in the GA

In order to evaluate which mechanisms are the most important for the CO2 splitting in the GA plasma, and how they can eventually be further improved, we investigated the dominant reaction pathways for the formation and loss of CO2 for the same conditions as in Fig. 4. The reactions are listed in Table 3 and their relative contributions to the overall CO2 loss and formation are presented in Fig. 6. This kinetic analysis was performed by looking at the time and volume integrated rates of the various processes for a complete gliding cycle of 10 ms. In the supporting information, we also plot the temporal evolution of the most important loss and formation rates of CO2, obtained by integrating the reaction rates over the entire reactor (see Fig. S4).

The most important process for CO2 loss is the dissociation of vibrationally excited states of CO2 upon collision with O atoms (L2v) with a relative contribution of about 80%. The same process, but upon collision of ground state CO2 with O atoms (L2g) has a relative contribution of 9.2%. Furthermore, the dissociation of vibrationally excited states of CO2 upon collision with any neutral species (M) also contributes for 7.3% (L3v). The relative contribution of the same process, but starting from ground state CO2, is only 0.21% (L3g). Besides, electron impact dissociation from the CO2 vibrational levels (L1v) and from the CO2 ground state (L1g) contribute for 2.6% and 0.70%, respectively. Compared with the electron impact dissociation reactions, the neutral reactions upon collision with O atoms have a lower energy.

<table>
<thead>
<tr>
<th>Process</th>
<th>Loss reaction</th>
<th>Process</th>
<th>Formation reaction</th>
</tr>
</thead>
<tbody>
<tr>
<td>L1v</td>
<td>$e + CO_2(v) \rightarrow e + CO + O$</td>
<td>F1</td>
<td>$CO + O_2 \rightarrow CO_2 + O^*$</td>
</tr>
<tr>
<td>L1g</td>
<td>$e + CO_2(g) \rightarrow e + CO + O$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>L2v</td>
<td>$CO_2(g) + O \rightarrow CO + O_2$</td>
<td>F2</td>
<td>$CO + O + M \rightarrow CO_2 + M$</td>
</tr>
<tr>
<td>L2g</td>
<td>$CO_2(g) + O \rightarrow CO + O_2$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>L3v</td>
<td>$CO_2(v) + O + M \rightarrow CO + O_2$</td>
<td>F3</td>
<td>$CO + O_2 \rightarrow e + CO_2$</td>
</tr>
<tr>
<td>L3g</td>
<td>$CO_2(g) + O + M \rightarrow CO + O_2$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>L4v</td>
<td>$CO_2(v) + O_2 + M \rightarrow CO_2 + 3M$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>L4g</td>
<td>$CO_2(g) + O_2 + M \rightarrow CO_2 + 3M$</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*a O2 represents the sum of the ground state and the vibrational states of molecular oxygen.*
requirement [19] and hence are more energy efficient. Note that reactions L2v and L2g are actually follow-up reactions of reactions L1v and L1g, as the O atom that reacts in reactions L2v and L2g is the result of CO2 splitting, either by reactions L1v and L1g, or reactions L3v and L3g. Nevertheless, once the first O atoms are formed upon CO2 splitting, the reactions L2v and L2g can occur in parallel to these other reactions, and thus we can consider them separately in this analysis.

Our calculation results reveal that the CO2 dissociation mainly proceeds from the vibrationally excited levels of CO2. The latter provide more energy efficient dissociation, because the vibrational energy can help overcome the activation energy barrier of the reaction and thus increase the reaction rate constant [26,27]. This is consistent with experimental investigations in literature. Indeed, experimental work for both a diverging electrodes gliding arc reactor [5] and a gliding arc plasmotron [6] shows that the presence of a very small quantity of water added into CO2 greatly reduces the power efficiency compared with pure CO2 at atmospheric pressure. This is explained by the fact that water can significantly reduce the vibrational excitation of CO2 molecules, because the energy is absorbed and quickly lost by water. Based on this, Nunnally et al. [6] concluded that non-equilibrium vibrational excitation plays the major role during CO2 dissociation in a gliding arc.

Additionally, there exist measurements in the literature, demonstrating that the vibrational temperature in the gliding arc is higher than the gas temperature, even at atmospheric pressure, although we cannot validate our model by direct comparison, as experimental data for the vibrational temperature in pure CO2 in classical gliding arc reactors do not yet exist. However, in a non-equilibrium gliding arc “tornado” discharge using CO2 doped with 1% N2 at a flow rate of 10 lpm and a power of 200 W, Nunnally et al. [49] estimated the vibrational temperature to be approximately 6000 K at atmospheric pressure, by comparing the theoretical and experimentally measured spectra for the N2 system, and this value is much higher than the reported rotational gas temperature of 2700 K ± 50 K. Therefore, these experimental results support our modelling results.

Some of the reactions plotted in Fig. 6(a) also occur in the opposite direction, hence, besides dissociation of CO2, the recombination of CO with O2, O and O− ions also takes place in the GA, giving rise to the formation of CO2 again and yielding a lower net conversion of CO2. The recombination reaction of CO with O2 molecules (F1, i.e., the opposite of L2) is the predominant production process of CO2, with a relative contribution to the overall CO2 formation amounting to 94%. The recombination reaction of CO with O atoms (F2, i.e., the opposite of L3) has a relative contribution of 5.2%, while the recombination of CO with O− ions (F3) only contributes for 0.068%. Other reactions play a negligible role towards CO2 formation (< 0.05%).

Note that the reverse reactions, especially the recombination of CO with O2 molecules, have only slightly lower rates than the rates of the most important loss processes, as depicted in Fig. S4 in the supporting information. Therefore, these reactions have a detrimental effect on the overall CO2 conversion. Indeed, when the rates of these reactions would become larger, they would inhibit further CO2 dissociation. This happens when a considerable fraction of the CO2 molecules is already converted into CO and O/O2, and especially at high gas temperature in the arc. When comparing the total loss of CO2, integrated over the entire arc and the whole gliding cycle, with the total formation of CO2, we obtain values of 3.8 x 1018 vs 3.5 x 1018 at the conditions under study. Thus, it is clear that about 92% of the CO2 converted in the GA, will be formed again, so the net conversion of CO2 into CO is much smaller than the initial loss of CO2. Therefore, the recombination of CO with O2 back into CO2 is clearly a limiting factor, which affects the further improvement of GA based CO2 conversion and its energy efficiency. This will be discussed in the next section.

4.5. How to improve the CO2 conversion and energy efficiency in the GA?

From previous section, we can clearly identify the limiting factors for energy efficient CO2 conversion in the GA. Therefore, in this section, we will propose solutions on how to further improve the performance of the GA for energy efficient CO2 conversion. First we will discuss the role of the vibrational levels in energy efficient CO2 conversion. Subsequently, we will look in more detail at the recombination of CO with O2, which contributes mostly to the CO2 formation at the conditions under study. Finally, we will elaborate on some ways to increasing the velocity difference between the GA and the gas flow, which can increase the fraction of CO2 that can be processed by the arc, and hence improve the conversion.

4.5.1. Promoting the vibrational kinetics

It is clear that non-equilibrium vibrational excitation of CO2 promotes energy efficient dissociation in the GA. This is also consistent with experimental investigations in literature [6]. Our results indicate that the population of the symmetric mode levels and the lower asymmetric stretch mode levels is much higher than that of the higher asymmetric mode levels. Therefore, these lower vibrationally excited levels mostly account for the total CO2 conversion, although there is still some overpopulation for the higher levels. The reason why especially the lower vibrational levels contribute to the CO2 conversion is because the vibrational energy distribution function tends to become more thermalized at high gas temperature [72]. Indeed, the energy exchange upon collision between vibrational levels and ground state molecules, which depopulates the vibrational levels, i.e., so-called VT relaxation, increases with gas temperature. Therefore, we should look for ways of inhibiting the VT relaxation process to increase the degree of overpopulation of the higher asymmetric mode levels.

A recent kinetic modelling of microwave plasma based CO2 conversion has shown that lower pressures, lower gas temperature and higher power densities (at least for pressures below 300 mbar) lead to more vibrational excitation, which is beneficial for the conversion [72]. However, our GA operates at atmospheric pressure, which is more convenient for industrial applications, so the solutions of reducing the gas pressure and increasing the power density (which only has beneficial effect at a pressure below 300 mbar [72]) are not practical. Therefore, we believe that the gas temperature should be reduced, to inhibit the VT relaxation, and thus to promote the role of the higher vibrational levels, and hence the conversion and energy efficiency. In this respect, enhancing the mixing between the GA and the cold gas can help to realize this goal, which was clearly indicated by our previous modelling for a 1D gliding arc [41] and by experimental work [6]. Furthermore, reducing the gas temperature will also result in a lowering of the recombination reactions, thus also improving the overall CO2 conversion (see next section). On the other hand, it will also lead to a drop in the dissociation rate constants by neutral particle collisions, and this has a detrimental effect on the conversion. Therefore, an optimized gas temperature should exist for GA based CO2 conversion, where the beneficial effect of a lower temperature, due to (i) a more pronounced non-equilibrium population of the highly excited vibrational levels, and (ii) lower recombination rates of CO back into CO2, exceeds the detrimental effect by the lower dissociation rate constants of dissociation upon collision with neutral particles. Finding out this optimal temperature is, however, not so straightforward with our 2D model, as the latter self-consistently calculates the gas temperature and it is not an input in the model. For this purpose, a 0D model, where the gas temperature can be introduced as an input parameter, could be more suitable [40].

Besides, because electron impact vibrational excitation of CO2 is mainly important for reduced electric field values (i.e., ratio of electric field over gas density) below 80 Td [71] (where 1 Td = 10^{-21} V/m^2), we should target to actively tune the reduced electric field to these values, by optimizing the reactor electrical operating parameters.
Finally, increasing the electron number density will also promote the vibrational excitation and thus selectively deliver energy to this most energy efficient CO2 dissociation pathway. It has been reported in literature [65] that adding noble gases, such as argon, to CO2 would improve the CO2 conversion and energy efficiency by increasing the electron number density, because argon has a lower breakdown voltage than CO2.

4.5.2. Reducing the recombination of CO with O2

It is clear from section 4.4 that the recombination reaction (F1), i.e., CO + O2 → CO2 + O, is mainly limiting the CO2 conversion and energy efficiency. In our model, we adopted the rate coefficient as proposed by Fridman [19]. However, to evaluate the effect of this recombination reaction on the overall CO2 conversion, we have performed some further simulations in which (i) we reduced the rate coefficient of this reaction by 50%, and (ii) we completely removed this recombination reaction from the model, as indicated in the legend of Fig. 7.

It is obvious from Fig. 7 that a lower rate coefficient of the recombination reaction yields a higher net CO2 loss rate. The CO concentration within the GA channel, and hence the influence of the recombination reaction on the CO2 formation, is minor till t = 1.7 ms. As a result, the different rate coefficients have a negligible effect on the net loss rate of CO2 up to 1.7 ms. Upon increasing CO concentration, the different rate coefficients do cause some deviation in the calculated net loss rates of CO2. After t = 7.5 ms, the formation rate of CO2 is even larger than the loss rate for k1 and k2, leading to a negative value of the net CO2 splitting rate. Of course, integrated over the entire GA cycle, the overall CO2 loss (or conversion) rate is still positive, but it is greatly reduced due to this important backward (recombination) reaction.

Fig. 8 shows the conversion and energy efficiency, calculated with the original rate coefficient (k1) [19], in comparison with the results obtained when this rate coefficient is divided by 2 (k2), as well as when the recombination reaction is removed from the model (k3). The conversion and energy efficiency increase only slightly when the recombination rate coefficient is divided by 2, while they rise from 2.8% to 4.0%, and from 33% to 47%, respectively, by removing the recombination reaction (CO + O2 → CO2 + O) from the model. Although the conversion is still low, the energy efficiency rises significantly. This clearly indicates that reducing the recombination of CO with O2 is quite promising to enhance the CO2 conversion and (especially) the energy efficiency.

To achieve this objective, we suggest to apply possible scavengers, catalysts or separation membranes, in order to remove the O2 molecules [31]. These are only suggestions, and they should of course be experimentally explored to evaluate the possibilities. On the other hand, the combination of a solid oxide electrolyser cell with a plasma set-up was already illustrated in [73] to be beneficial for the CO2 conversion, and it works according to the same principle. In this way, the local concentration of O2 molecules within the arc channel, and hence the net formation of CO2 by the recombination reaction (CO + O2 → CO2 + O), could be reduced, because there is not enough reactant (O2) available for the backward reaction from CO into CO2 (F1).

H2 or CH4 could act as possible scavengers for atomic oxygen, forming H2O. This possibility was already illustrated to be beneficial for O trapping in literature, based on a combined plasma chemical kinetics model and experiments for CO2 conversion in another type of plasma [74]. The trapping of O atoms might be able to promote the CO2 conversion by (i) inhibiting the recombination reaction F2 [40], and (ii) by avoiding the formation of O2, which will inhibit the recombination reaction F1. Experiments in literature have indeed revealed that the addition of H2 or CH4 in a GA reactor can improve the conversion of CO2 [6,54], but the enhanced conversion of CO2 cannot be simply, or entirely, attributed to the inhibited recombination reactions. This is because the H atoms or CHx radicals produced by H2 or CH4 dissociation can also contribute to CO2 dissociation. Moreover, the removal of O atoms will also inhibit the dominant mechanism of CO2 splitting, i.e. the dissociation of CO2 upon collision with O atoms (L2v, L2g) and thus it might also exhibit a negative effect on further improving the CO2 conversion. Therefore, the reason why adding H2 or CH4 promotes the CO2 conversion is not necessarily attributed to their scavenging role in consuming the O atoms. Indeed, the direct involvement in CO2 splitting...
by the reversed water gas shift reaction \((\text{CO}_2 + \text{H}_2 \rightarrow \text{CO} + \text{H}_2\text{O})\) has been verified to be a very important path for CO splitting into CO when CH\(_4\) [54] or H\(_2\) [6] is added into a CO\(_2\) GA plasma. Moreover, the addition of H\(_2\) or CH\(_4\) can increase the electron density by inhibiting electron attachment to O\(_2\) (which is an electron-negative gas), and this can also contribute to a higher CO\(_2\) conversion.

The idea of using a catalyst with a high surface interaction for O atoms to recombine into O\(_2\) [75] or for O\(_2\) adsorption is probably not very effective, because the O\(_2\) molecules would be released back to the plasma phase and again undergo recombination with CO. In contrast, a more advanced catalytic process would be an alternative form of chemical looping, in which the O or O\(_2\) is captured in the plasma set-up and then used as oxidizing agent in a second set-up [76,77]. However, this is only a concept, and has not been demonstrated yet for a GA reactor.

The third method, based on separation membrane technology, would transport the O\(_2\) molecules (or O atoms) away from the reaction mixture. For example, by combination of a solid oxide electrolyser cell with a plasma set-up, Tagawa et al. [73] and Mori et al. [78,79] have observed an increasing CO\(_2\) conversion by placing an O\(_2\) trapping membrane into a CO\(_2\)/CH\(_4\) or CO\(_2\) discharge, in order to separate O\(_2\) from the reaction mixture.

Besides the effect of possible scavengers, catalysts or membranes to remove the oxygen, as mentioned above, we believe that the recombination of O\(_2\) could also be avoided or minimized by providing effective quenching of the high temperature in the arc zone, due to mixing with cold gas at very fast cooling rates. This could be especially beneficial in the relaxation stage of the GA (around 8 ms) when the discharge current is low, and the CO\(_2\) loss rate is minor, but the recombination rate of CO with O\(_2\) is still very large due to the very high gas temperature, leading to net CO\(_2\) formation. Indeed, an effective quenching of the residual plasma temperature can help to decrease the recombination reaction rate and inhibit the CO\(_2\) formation in this stage, leading to an improved conversion and energy efficiency. We believe that such a quenching of the plasma temperature could be realized by improving the reactor geometry and/or optimizing the flow conditions, but further studies are needed to elaborate on these solutions.

4.5.3. Increasing the CO\(_2\) fraction to be treated by the arc due to a velocity difference between GA and gas flow

Besides promoting the vibrational kinetics and reducing the recombination reaction of CO into CO\(_2\), another way to improve the CO\(_2\) conversion would be to enhance the CO\(_2\) fraction to be treated by the arc, by better mixing of the GA and the cold gas flow. This can be realized when there is a velocity difference between the GA and the gas flow. Several experimental studies indeed have shown that the arc gliding velocity can be slightly lower than the gas velocity [14,33]. We present here some simulation results, showing that there can indeed be a (small) difference between the arc and gas flow velocity. We can distinguish two different ways to realize this.

1) Smooth velocity difference due to the arc bending

The first possible reason for a lower arc velocity vs gas flow velocity is related to the arc bending, and therefore the existence of zones with incresed electric field outside the arc centre. The latter indeed leads to a separation of the arc centre (with the maximum electron number density) and the position with maximum reduced electric field, as presented in Fig. 9. This is caused by the fact that in the symmetry plane, when the arc is highly bended, some parts of the arc in the downstream region of the arc centre are positioned closer to each other. This increases the electric field strength in this region and causes a gradual ionization of the gas in the downstream region. The latter will result in a slightly lower arc velocity compared to the gas velocity. Likewise, near the walls (cathode and anode), the maximum reduced electric field, and hence the gradual ionization, appears in the upstream region of the arc centre, which results in a slightly higher arc velocity than the gas velocity. Thus, the GA moves a bit slower than the gas flow in the central part of the reactor and a bit faster in the regions near the walls. At t = 2.5 ms, our calculation predict a GA velocity of 5.9 m/s in the centre, compared to a gas flow velocity of 7.4 m/s. The ratio of gas velocity to arc velocity is thus 1.2, which is in reasonable agreement with experiments [14,33].

We have also performed calculations at higher gas flow velocity, and the results show that this leads to an increased velocity difference between the arc and gas flow. For example, with the same gas flow rate of 2.5 L/min, assuming the flow passing through a channel with depth of 1 mm, which is only half of the value in our standard model, the gas flow velocity at the same time instant t = 2.5 ms was calculated to be 8.3 m/s, with a GA velocity of 5.8 m/s, thus yielding a ratio of gas velocity vs arc velocity of 1.4. This clearly shows that the velocity difference between GA and gas flow will be higher for higher gas flow velocities, which is also reported in experiments [14,33]. Correspondingly, our calculated conversion increases from 2.78% to 4.4%, although the energy efficiency only increases from 32.8% to 34%. Although this is an artificial method, we can show in this way that the treatment capacity can be enlarged by increasing the local gas velocity and hence the relative velocity between gas flow and GA. Increasing the local gas velocity can be realized by modifying the reactor setup and hence the flow configuration at a fixed gas flow rate, for example by shortening the narrowest gap separation of both electrodes [40] or by reducing the distance between the nozzle exit and the reactor [80] or by decreasing the nozzle internal diameter [81]. Indeed, following such methods, increased conversions were reached experimentally [40,80,81]. However, we should also mention that simply adjusting these parameters is not a proper way to enhance the treatment capacity of the GA reactor, because it might give rise to an extreme increase in the gas velocity, which may greatly reduce the effective residence time of CO\(_2\) in the GA volume. This is of course detrimental for the CO\(_2\) conversion. Moreover, the high gas velocity will bring a strong cooling effect and hence a lower gas temperature; the latter can be beneficial (to promote the vibrational kinetics and/or reduce the recombination reactions), but it may also be detrimental (due to the reduced dissociation reaction rate constant), as we discussed in section 4.5.1 above. Therefore, the above mentioned operating parameters should be optimized in a suitable range, to guarantee an improvement in conversion and energy efficiency [82].
Sudden velocity difference due to back-breakdown events

Besides the smooth reduction in GA velocity explained above, another reason for the lower arc velocity vs gas flow velocity is related to the instabilities of the arc and to secondary breakdows, also called back-breakdown, causing a reduction in arc length [83].

The back-breakdown phenomena, which result in a fast shortening of the arc as a result of breakdown between different parts of the arc (instead of between the electrodes) often take place in a GA, especially at higher gas flow rates, as also mentioned in section 4.1. These shortcuts effectively appear as a lag of the arc velocity compared to the gas flow and could be an efficient mechanism for the treatment of a larger gas fraction. As explained at the end of section 4.1, this effect is not taken into account in previous sections, because at the gas flow rate of 2.5 L/min, our high speed camera did not record any back-breakdown events.

Fig. 10 illustrates a back-breakdown event, recorded by the high speed camera at a flow rate of 5 L/min. Indeed, at a high gas flow rate (above 2.5 L/min), the GA discharge is unstable and it has a rather irregular shape. When some parts of the GA (see points A and B in Fig. 10) get closer to each other, the electric field there increases. Once the potential difference between these two parts, and hence the local electric field, exceeds the critical breakdown electric field [84], a new discharge channel is established (see middle panel) and the old discharge channel disappears very fast. This causes a drop in the GA velocity as compared to the gas flow velocity.

Although several experiments [34,83] have been performed to study the back-breakdown events, it is not straightforward to establish a self-consistent back-breakdown model, since this behaviour is mostly stochastic by nature and the arc instabilities are not well defined. To investigate here the influence of the back-breakdown events on the CO2 conversion, we have initiated this process by establishing an artificial plasma channel, which is triggered on a regular or irregular basis with respect to the arc path or time, i.e., after every certain distance or period. Details on how the back-breakdown model is established can be found in [38], as well as in the supporting information of our paper.

Fig. 11 illustrates the effect of the back-breakdown events on the calculated conversion and energy efficiency. The power needed to initiate the back-breakdown events is included in the determination of the total plasma power and hence in the SEI value in Eq. (3), as well as the calculation of the energy efficiency in Eq. (4) (see section 2). It is clear that the back-breakdown events yield an improved CO2 conversion and energy efficiency, compared with the case without back-breakdown, because a larger fraction of CO2 is treated by the newly established discharge channel. This also explains why a larger number of back-breakdown events can enhance the CO2 conversion and energy efficiency (see cases 2, 3, 4 and 5). Moreover, more back-breakdown events also result in a lower overall gas temperature, as is clear from Fig. S7 of the supporting information, because the heat is now spread over a larger domain and not only within the initial arc channel. This lower gas temperature can have beneficial or detrimental effects on the overall CO2 conversion, as explained above.

As discussed above, the occurrence of the back-breakdown events is closely linked with two factors, i.e. the arc instabilities and a sufficiently high arc voltage drop. The former leads to a rather irregular arc shape and a non-stable discharge, increasing the probability of a closer interaction between two separated parts of the GA. The latter can
ensure a high enough electric field between the two separated parts of the arc, to ignite a new discharge channel. In order to satisfy these two essential requirements, besides increasing the gas flow rate, the gas flow velocity must also be increased by modifying the reactor setup and hence the flow configuration under a fixed gas flow rate, as discussed above.

4.5.4. Summary of the proposed improvements

Finally, in Fig. 12, we schematically summarize the improvement in the CO2 conversion and energy efficiency, as proposed and predicted by our model. The CO2 conversion and energy efficiency are about 2.78% and 32.8% (calculated) or 2.90% and 34.3% (measured) at the standard conditions investigated, i.e., a gas flow rate of 2.5 L/min and a plasma power of 40 W, corresponding to a SEI of 0.25 eV/molecule. However, these values can be improved according to the model predictions, up to a conversion of nearly 4% and a corresponding energy efficiency of 47% (see point b) by inhibiting the recombination reaction of CO with O2. Furthermore, if the gas fraction that can pass through the arc zone could be enhanced, for instance by modifying the reactor setup and hence the flow configuration to realize a higher relative velocity between arc and gas flow, the conversion and energy efficiency are predicted to increase to 4.4% and 34%, respectively (see point g). Finally, the occurrence of back-breakdown events, which induce an abrupt decrease in gas flow velocity and GA velocity, in case of a gas flow rate of 5 L/min (where the back-breakdown events indeed can take place), can also help to increase the conversion, although the effect seems to be rather limited, with a maximum conversion up to 2.6%, while the energy efficiency would increase up to 41% (see point f).

The proposed solutions yield some improvement in conversion and energy efficiency, but these model predictions still need to be verified by experiments. We hope that our model predictions will inspire experimental researchers to try out these modifications. Furthermore, the improvements are probably still too limited for industrial application of the GA for CO2 conversion. Indeed, although the energy efficiency is quite good, the conversion is still very limited. Hence, more drastic modifications would be needed, e.g., in the gas flow pattern or the source design, to significantly increase the fraction of gas that can pass through the arc. One possible suggestion would be the reverse vortex flow gliding arc, which is based on cylindrical electrodes, and which allows a larger fraction of the gas to pass through the arc, yielding higher CO2 conversions, as demonstrated by [6,50,51].

5. Conclusions

In this work we studied the CO2 conversion in a GA plasma, by means of a combined experimental and 2D modelling approach. We compared our measured and calculated CO2 conversion and corresponding energy efficiency, as well as the electron number density in the arc, and obtained reasonable agreement. This indicates that our model can provide a realistic picture of the plasma chemistry and can be used to elucidate the underlying mechanisms and the dominant reaction pathways for the GA based CO2 conversion.

We presented the typical arc plasma characteristics, such as the electron number density, electron temperature and gas temperature, as well the CO molar fraction, for one entire arc gliding cycle, as calculated by our model. These results clearly show that the GA plasma has a strong non-equilibrium character, because the electron temperature is much higher than the gas temperature, and the highly energetic electrons can induce several different chemical reactions. This explains the better performance of the GA for CO2 conversion, yielding a much higher energy efficiency for a fixed value of the conversion, than pure thermal conversion, for which the energy is distributed over all degrees of freedom, including those not effective for the CO2 conversion.

We also performed a chemical kinetics analysis of the modelling results, which enables us to identify the important species and reactions playing a role in the CO2 splitting, i.e., the main production and loss pathways of CO2. This allows us to gain sufficient insight into the entire process, and to identify the limiting factors for CO2 conversion, and thus to propose solutions for improving the CO2 conversion. Our model predicts that the most important process for CO2 conversion is the dissociation of vibrationally excited states of CO2 upon collision with O atoms, indicating that the CO2 vibrational levels significantly contribute to the CO2 dissociation. This can explain the good energy efficiency of CO2 conversion in a GA plasma, as compared to some other plasma types.

We believe that, when it is possible to actively tune the reduced electric field (i.e., E/n ratio) in the plasma, by optimizing the reactor electrical operating parameters, or when we can increase the electron number density, as well as inhibit the VT relaxation processes by decreasing the gas temperature, we should be able to further promote the vibrational excitation and selectively deliver energy to the CO2 dissociation via this energy efficient pathway. This should lead to some further improvement in the energy efficiency of CO2 conversion in the GA.

Furthermore, our calculation shows that the reverse reactions, especially the recombination of CO with O2 molecules (and to a lower extent with O atoms), have a non-negligible rate, compared to the CO2 loss rate. Therefore, these reactions have a detrimental effect on the overall CO2 conversion. Thus, in order to further improve the CO2 conversion, the reversion reactions should be inhibited or at least reduced. We clearly demonstrate this by running the model with different reaction rate coefficients for recombination, and when this recombination reaction is entirely removed, the calculated CO2 conversion and energy efficiency rise from 2.8% and 33%, to 4.0% and 47%, respectively.

Finally, our simulation shows that the molar fraction of CO2 within the arc centre is very low, indicating that the local CO2 conversion is nearly complete, but because the fraction of treated CO2 within the arc is very limited, the overall CO2 conversion is also limited. Therefore, we should look for ways to increase the CO2 fraction to be treated by the arc, in order to further improve the GA based CO2 conversion. Increasing this treated gas fraction can be realized when there is a velocity difference between the GA and the gas flow, so that new fractions of the CO2 gas can pass through the arc, while the converted fraction (i.e., CO, O and O2) will leave the active arc region, before it.
can recombine back into CO2. We therefore discuss possible ways of increasing the relative velocity between GA and gas flow. The first way to realize this is by increasing the local gas velocity without changing the gas flow rate, for instance by modifying the reactor setup and hence the flow configuration. Indeed, at a high gas velocity, there is a larger difference between GA and gas flow velocity, so we also investigated the effect of these back-breakdown events on the calculated CO2 conversion and energy efficiency. Our calculations clearly indicate that the back-breakdown events, which generally take place at a high gas flow rate, can help to further increase the CO2 conversion and energy efficiency.

This study is of great interest for GA based CO2 conversion, as we were able to elucidate the main underlying mechanisms and chemical reactions of the conversion process by means of a model that was validated by experiments. In general, we illustrated that GA based CO2 conversion is quite promising, when compared with the classical thermal CO2 conversion process, as well as with other plasma types. This is attributed to its non-equilibrium character, promoting the vibrational kinetics. However, we believe there is still room for improvement. Indeed, we could identify the limiting factors of the CO2 conversion in the GA, and thus propose solutions on how to further improve the performance.

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Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/jcej.2017.07.133.

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