

This item is the archived peer-reviewed author-version of:

DBD in burst mode : solution for more efficient CO_2 conversion?

Reference:

Ozkan Alp, Dufour T., Silva T., Britun N., Snyders R., Reniers F., Bogaerts Annemie.- DBD in burst mode : solution for more efficient CO_2 conversion?

Plasma sources science and technology / Institute of Physics - ISSN 0963-0252 - 25:5(2016), 055005

Full text (Publishers DOI): <http://dx.doi.org/doi:10.1088/0963-0252/25/5/055005>

DBD in burst mode: solution for more efficient CO₂ conversion?

A. Ozkan^{1,2}, T. Dufour¹, T. Silva³, N. Britun³, R. Snyders³, F. Reniers¹ and A. Bogaerts²

¹ *Université Libre de Bruxelles, Chimie analytique et chimie des interfaces (CHANI), Campus de la Plaine, Bâtiment A, CP255, boulevard du Triomphe, 1050 Bruxelles, Belgium*

² *Universiteit Antwerpen, Research group PLASMANT, Universiteitsplein 1, 2610 Antwerpen-Wilrijk, Belgium*

³ *Université de Mons, Chimie des Interactions Plasma-Surface (ChIPS), CIRMAP, 23 Place du Parc, 7000 Mons, Belgium*

Keywords: CO₂ conversion; cold atmospheric plasma; flowing DBD; burst mode; filamentary discharge; electrical characterization.

Abstract

The CO₂ conversion into value-added products has gained significant interest over the last years as the greenhouse gas concentrations constantly increase due to anthropogenic activities. We report here on experiments for CO₂ conversion by means of cold atmospheric plasma using a cylindrical flowing dielectric barrier discharge (DBD) reactor. A detailed comparison of this DBD ignited in so-called burst mode (i.e., where an AC voltage is applied during a limited amount of time) and pure AC mode is carried out to evaluate their effect on the conversion of CO₂ as well as on the energy efficiency. Decreasing the duty cycle in the burst mode from 100% (i.e., corresponding to pure AC mode) to 40% leads to a rise in the conversion from 16 to 26% and to a rise in the energy efficiency from 15 to 23%. Based on a detailed electrical analysis, we show that the conversion is correlated with the features of the microfilaments. Moreover, the root-mean-square voltage in the burst mode remains constant as a function of the process time for duty cycles < 70%, while a higher duty cycle or the usual pure AC mode leads to a clear voltage decay by more than 500 V over approximately 90 seconds, before reaching a steady state regime. The higher plasma voltage in burst mode yields a higher electric field, hence increasing the electron energy, and thus their involvement into the CO₂ dissociation process, which is an additional explanation for the higher CO₂ conversion and energy efficiency in the burst mode.

Introduction

New records in greenhouse gas concentrations are reported every year. Especially carbon dioxide (CO₂) is considered as an inevitable reaction product of human activity. Indeed, the anthropogenic activities have led to increasing atmospheric CO₂ concentrations over the last 150 years, from 280 to almost 400 ppm in 2014 [1]. This means higher global temperatures and more extreme weather phenomena such as heat waves, floods, ice melting, rising of sea

levels and ocean acidity [2-7]. This concern has recently been reported during the three weeks COP-21 conference in Paris to prompt governments into taking strong measures in order to limit global warming. Nowadays, the political authorities want to limit the production of CO₂ by promoting renewable energies, non-carbon energies, geological storage in deep underground [8-10] or by taxing the tons of CO₂ produced. As an innovative alternative, there is increasing interest in the possibility to reutilize CO₂, i.e. in a "cradle to cradle" approach, where waste of all types is being recycled. Carbon dioxide – the ultimate gaseous combustion product of carbon compounds – could be reutilized, although it is not straightforward, because it is a particularly stable molecule ($\Delta G_f \ll 0$).

In recent years, many publications report on CO₂ recycling following conventional organic reactions [11], electrochemical reduction in solution [12] or reduction in plasma phase [13]. The latter technique appears very promising, as it can directly operate in the gaseous phase while electrochemistry techniques must be carried out in liquid phase. Moreover, if it is mixed with hydrogen, water or methane, CO₂ can easily be reduced into syngas (CO/H₂) and small organic molecules [14-21]. The synthesis of value added compounds from CO₂ considered as a "raw material" is a very interesting concept for the industrial sector. However, as CO₂ corresponds to a particularly stable oxidized state of carbon, its reduction is not spontaneous and requires significant energy input. In cold atmospheric plasmas, such as dielectric barrier discharges (DBD), this energy is provided by electric power. The latter mainly heats the electrons, while the gas itself can remain near room temperature, thus allowing the reaction to proceed at mild reaction conditions (ambient pressure and temperature). Indeed, the energetic electrons will activate the gas molecules by excitation, ionization and dissociation reactions, creating reactive species that can easily form new molecules.

A DBD is commonly supplied with an AC voltage (pure AC mode) (e.g., [22-24]) and sometimes by high voltage nano or micro pulses [25-35] (pulsing mode). According to Jiang et al. [25], supplying a DBD with high voltage nanopulses enhances its stability over time, as well as its uniformity, since no irregular distribution of microdischarges is revealed. Such a power delivery presents great benefits for plasma polymerization, thin layer deposition and surface treatments since no significant texturization appears and it allows for instance an optimal deposition of a subsequent barrier layer [36-38]. This configuration is also of interest for gas treatment since a larger number of microdischarges is produced, increasing the probability for a single gas molecule to pass through the discharge and interact with a single microdischarge. Ndong et al. explain that positive voltage pulses create discrete discharge channels uniformly arranged on the dielectric surface in a streamer-like regime, while negative voltage pulses mostly tend to generate a uniform discharge along the active electrode such as a glow-like discharge [39, 40]. Song et al. investigated a bipolar pulsed discharge for the CO₂ reforming of methane where both the CO₂ and CH₄ conversion are higher when applying a pulse mode compared to a classical sinusoidal AC mode [41]. Some other authors have also compared a unipolar and bipolar DC discharge [42-44] and showed that the CO₂ conversion slightly increases upon rising pulse frequency.

Besides the usual pure AC and pulsing modes, a DBD can also be supplied by an AC voltage that is switched off at regular times, at a frequency much lower than the AC frequency. This so-called burst mode is well known in manufacturer's manuals and also in the literature dedicated to power supplies [45-53]. Benard et al. [45] have demonstrated how the production of small perturbations to the classical alternative voltage (i.e. burst mode) can enhance the stability of the DBD. However, to the authors' knowledge, there exist no papers yet in

literature dealing with cold atmospheric plasma sources operating in this so-called burst mode for gas conversion.

In the present article, we investigate the splitting of CO₂ using a flowing DBD operating in burst mode and we compare its performance with the usual pure AC mode. This burst mode corresponds to a sinusoidal voltage with a signal frequency of 28.6 kHz (period of 35.7 μs), which is switched on and off at a repetition frequency between 400 and 900 Hz. The influence of this mode is investigated on the CO₂ conversion, the energy efficiency and the microdischarge features (i.e., number, lifetime and charge of the microdischarges) and we compare the results with the same DBD that is supplied with a pure AC voltage (duty cycle = 100%). Electrical characterization, mass spectrometry and optical emission spectroscopy are applied to understand how and why the CO₂ conversion is enhanced.

Experimental set-up

DBD reactor and voltage measurements

The CO₂ gas flows through a cylindrical DBD reactor as shown in Figure 1. The CO₂ flow rate is set at 200 mL_n.min⁻¹ and the operating frequency is fixed at 28.6 kHz. The central cylindrical copper electrode has a diameter of 22 mm and a length of 120 mm. This inner electrode is powered by an AC high voltage, whereas the outer electrode is grounded. The latter is a mesh, made of stainless steel, with length of 100 mm (thus defining the discharge length), and placed around a dielectric barrier (2 mm in thickness), made of alumina and with inner diameter of 26 mm, yielding a discharge gap of 2 mm, and thus a discharge volume of 15.1 cm³. This corresponds to a residence time of the gas equal to 4.5 s for the given gas flow rate of 200 mL_n.min⁻¹. The applied power is provided by an AFS generator G10S-V and a transformer, which amplify the sinusoidal signal of the alternating voltage and which allow us to pulse the voltage in a controlled manner.

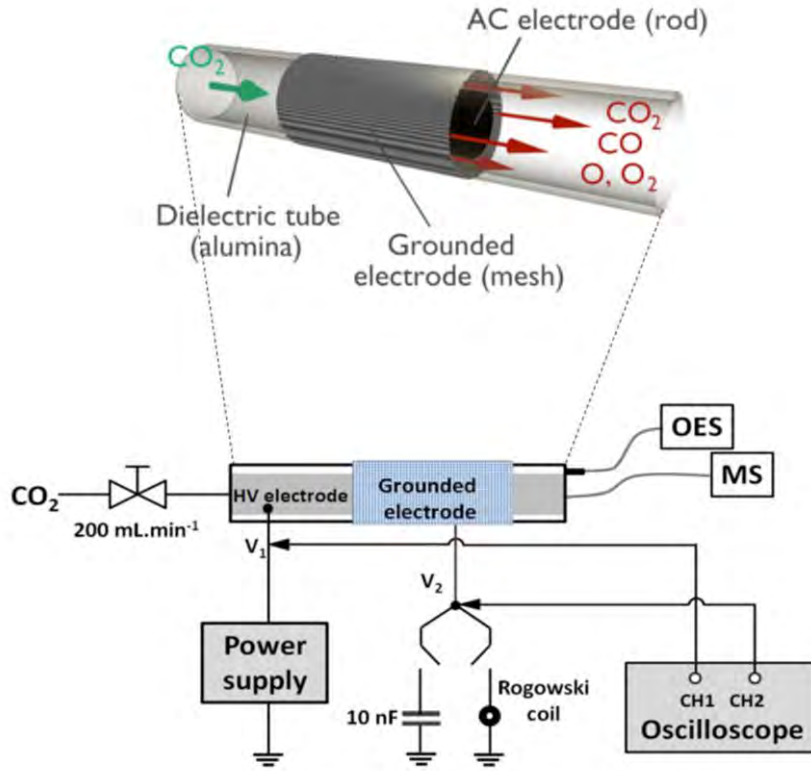


Figure 1 : Schematic diagram of the experimental set-up.

In the present study, the signal frequency remains fixed at $f_{\text{signal}}=28.6$ kHz (period of 35.7 μs). Furthermore, T_{ON} and T_{OFF} are defined as the time of plasma ignition and the time when no power is applied to the reactor. Except for the analysis of T_{ON} and T_{OFF} at constant duty cycle, the main results described in this article are related to the influence of the duty cycle. In this latter case, T_{ON} remains fixed at 1 ms while T_{OFF} is changed between 0 and 1.5 ms, corresponding to a duty cycle (D_{cycle}) between 100 and 40% . Therefore, the repetition frequency of the burst mode, $f_{\text{repetition}}=1/(T_{\text{ON}}+T_{\text{OFF}})$, is varied between 400 and 900 Hz. The voltage as a function of time is plotted in Figure 2 for high voltages delivered either in pure AC mode (a) and in burst mode (b). As the total power applied to the plasma (P_{applied}) is fixed at 50 W in all cases, the power during the plasma ignition (P_{plasma}) in burst mode is always higher than in pure AC mode, since $P_{\text{plasma}}(\text{W}) = P_{\text{applied}}(\text{W}) / D_{\text{cycle}}(\%)$.

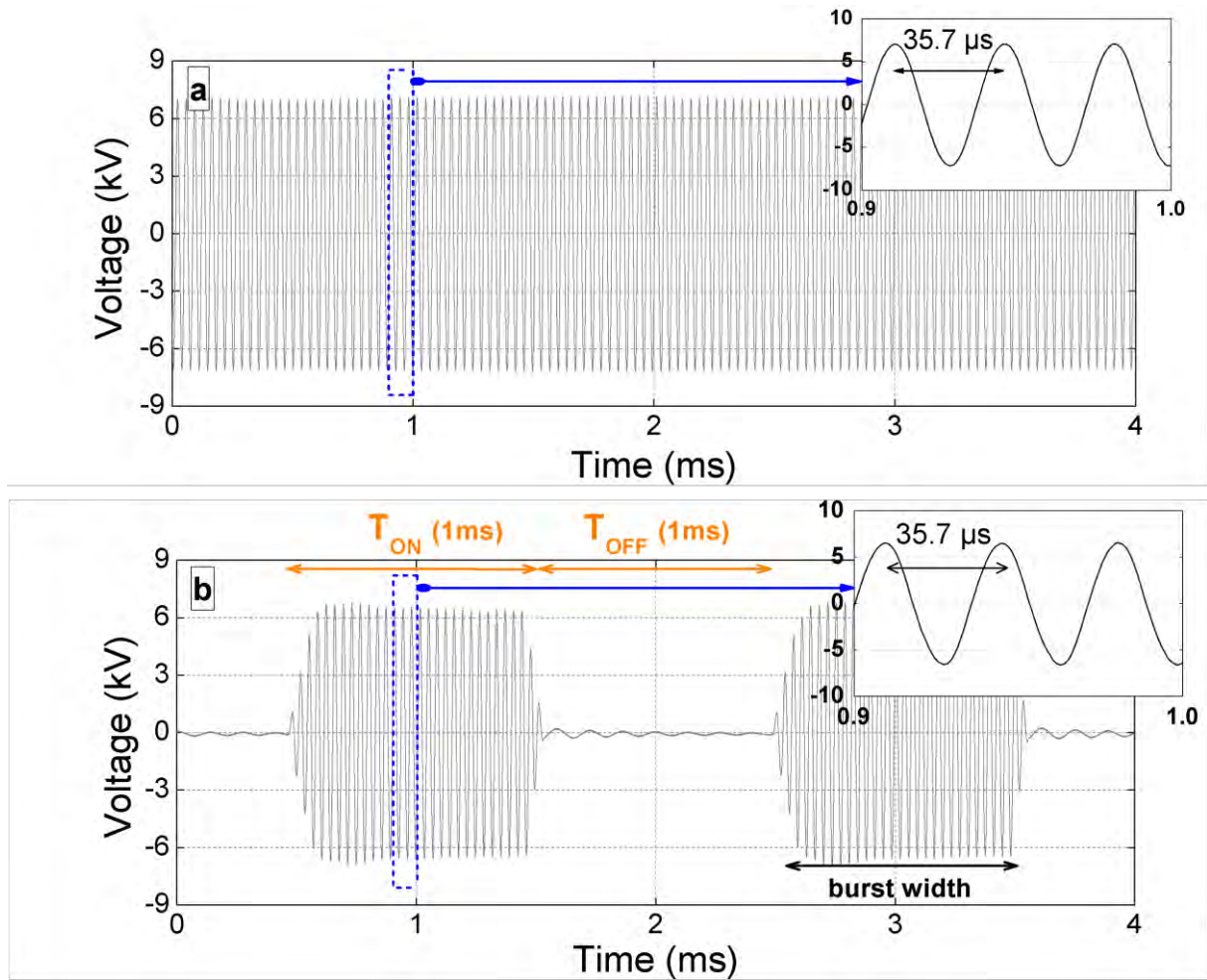


Figure 2 : High voltage signals applied to the discharge, with frequency of 28.6 kHz (or period of 35.7 μ s), in (a) pure AC mode, and (b) burst mode, with a duty cycle of 50% ($T_{ON} = T_{OFF} = 1$ ms).

Mass spectrometry (MS)

The gaseous products from the post-discharge are analyzed by a mass spectrometer operating at atmospheric pressure (Hiden analytical QGA). The electron energy in the ionization chamber is set at 70 eV and a secondary electron multiplier (SEM) is used as detector. The CO_2 conversion (χ_{CO_2}) is calculated according to the following equation, where I stands for the intensity of the signals:

$$\chi_{CO_2} (\%) = \frac{I_{CO_2 \text{ without plasma}} - I_{CO_2 \text{ with plasma}}}{I_{CO_2 \text{ without plasma}}} \times 100\%$$

The energy efficiency of the CO_2 conversion is calculated from χ_{CO_2} , the enthalpy of the splitting reaction ($CO_2 \rightarrow CO + \frac{1}{2} O_2$), namely $\Delta H_{298K}^0 = 279.8 \text{ kJ.mol}^{-1} = 2.9 \text{ eV.molecule}^{-1}$ and the specific energy input (SEI), according to the following equation:

$$\eta_{CO_2} (\%) = \chi_{CO_2} (\%) \cdot \frac{\Delta H_{298K}^0 (\text{eV.molecule}^{-1})}{SEI (\text{eV.molecule}^{-1})}$$

The SEI is defined as follows:

$$SEI (eV.molecule^{-1}) = \frac{P_{applied} (J.s^{-1})}{Gas\ flow\ rate (cm^3.s^{-1})} \times \frac{6.24 \times 10^{18} (eV.J^{-1}) \times 24500 (cm^3.mol^{-1})}{6.022 \times 10^{23} (molecule.mol^{-1})}$$

In principle, GC analysis is more accurate for determining the CO₂ conversion, but it was not available for this study. We believe that MS analysis is also sufficiently accurate at the current conditions.

Electrical measurements

A Tektronix DPO 3032 oscilloscope is used to perform electrical measurements (during T_{ON}). The first channel is dedicated to the monitoring of the high voltage (V₁) with a Tektronix P6015A probe while the second channel allows measuring the voltage passing through a capacitor, to obtain the power absorbed by the plasma via the Lissajous method [54-58], or through a Rogowski coil (Pearson 2877) to probe the current. The capacitor or the Rogowski coil is placed in series with the DBD. The obtained current profiles as a function of time, also called oscillograms, correspond to 2 different types of current: the plasma current and the dielectric current. These two can be simply distinguished as the dielectric component is a sinusoidal-like signal, whereas the plasma component corresponds to a hundred of microdischarges occurring at each half period, since an atmospheric pressure DBD operating in CO₂ generally operates in a filamentary regime [24, 59-61]. Analysis of these microdischarges is important to better understand the CO₂ splitting process. A statistical study will be carried out for several periods in order to evaluate the magnitude of different electrical parameters of these microdischarges. By applying a numerical method described in our previous article [61], the number of microdischarges (N_{md}), their lifetime (L_{md}), the plasma charge accumulation (Q_{pl}) and the resulting plasma current (i_{pl}) can be achieved. These data are collected for several periods and then averaged over a single period in order to have statistically valid results. In particular, N_{md} has been measured during T_{ON}=1ms and multiplied by D_{cycle} in order to consider the same residence time. Likewise, the applied voltage (V_{DBD}) can also be expressed as the sum of two voltages: the dielectric voltage (V_{diel}) and the plasma voltage (V_{pl}), which typically represents 80% of the total V_{DBD} [61, 62].

Optical emission spectroscopy (OES)

The OES measurements are performed with an Andor Shamrock-500i spectrometer including an Andor DU420A-OE CCD camera. Each spectrum is acquired for an exposure time of 5 s and 7 accumulations. Of particular interest is the CO emission band headed at 483.3 nm from the CO Angstrom system (B¹Σ⁺ → A¹Π; v'=0 → v''=1) shown in Figure 3. Its ro-vibrational structure is calculated and fitted to the experimental spectra in order to determine the rotational temperature (T_{rot}) and hence the gas temperature (T_{gas}), assuming T_{rot}≈T_{gas} [63-65]. For the sake of comparison, the gas temperature is also measured using another band: the first positive system (FPS) of N₂ is analyzed by adding 10% of nitrogen to CO₂. In this case, the temperature is measured via a line-ratio peak formula, specific to this system [66, 67], using the peaks at 775.3 and at 773.9 nm.

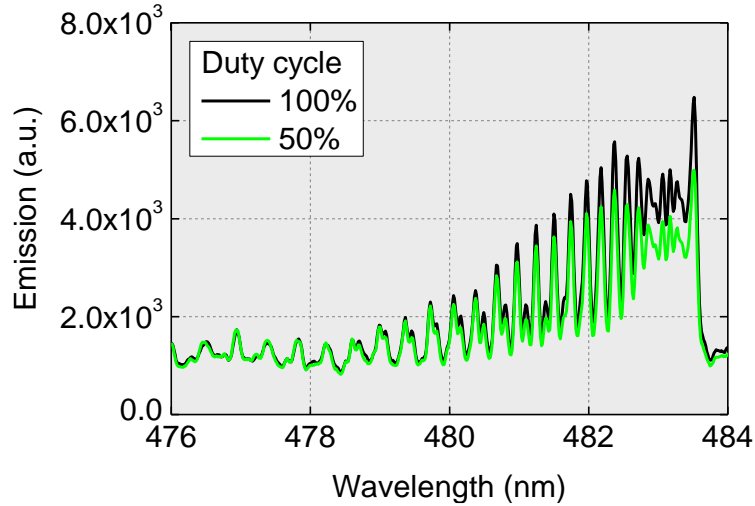


Figure 3 : CO band from the $[B^1\Sigma^+ \rightarrow A^1\Pi; v'=0 \rightarrow v''=1]$ Angstrom System in the DBD supplied with pure CO_2 , at two different duty cycles.

Results and Discussion

CO₂ conversion and energy efficiency

The CO_2 conversion and energy efficiency as a function of the duty cycle of the burst mode are depicted in Figure 4a. The conversion obtained in the pure AC mode ($D_{\text{cycle}}=100\%$) is 16.1%, which is significantly lower than in the burst mode at 40% duty cycle (i.e., 25.8%). Decreasing the duty cycle at constant applied power thus results in a higher CO_2 splitting. As evidenced in Figure 4b, smaller duty cycles yield higher peak powers, e.g. 100 W for $D_{\text{cycle}}=50\%$ versus 50 W for $D_{\text{cycle}}=100\%$. These higher peak powers delivered within shorter times can explain the higher CO_2 conversion. However, we will also show at the end of the paper that the enhancement of the burst mode is not only attributed to the higher peak power, but also to other effects (see below). The corresponding energy efficiency also increases upon decreasing duty cycle from 14.5% (at $D_{\text{cycle}}=100\%$) to 23.1% (at $D_{\text{cycle}}=40\%$).

On the other hand, keeping the duty cycle constant but increasing the burst width from 1 ms to even more than 20 ms has no effect on the CO_2 conversion and energy efficiency, as illustrated in Figure 5. Therefore, in the next sections we will only investigate the influence of T_{OFF} (or duty cycle or repetition frequency) on the electrical properties of the discharge, and more specifically on the microdischarge features. First we will show the effect of the duty cycle on the evolution of the applied voltage with increasing process time, to check the time-stability of the discharge. Next, the influence of the duty cycle on the current oscillograms and on the microdischarge properties, as well as on the gas temperature will be studied. Finally, the burst mode and the normal AC mode will be compared at the same delivered power, to illustrate that the enhancement of the burst mode is not only attributed to the higher peak power, but also due to the microdischarge properties.

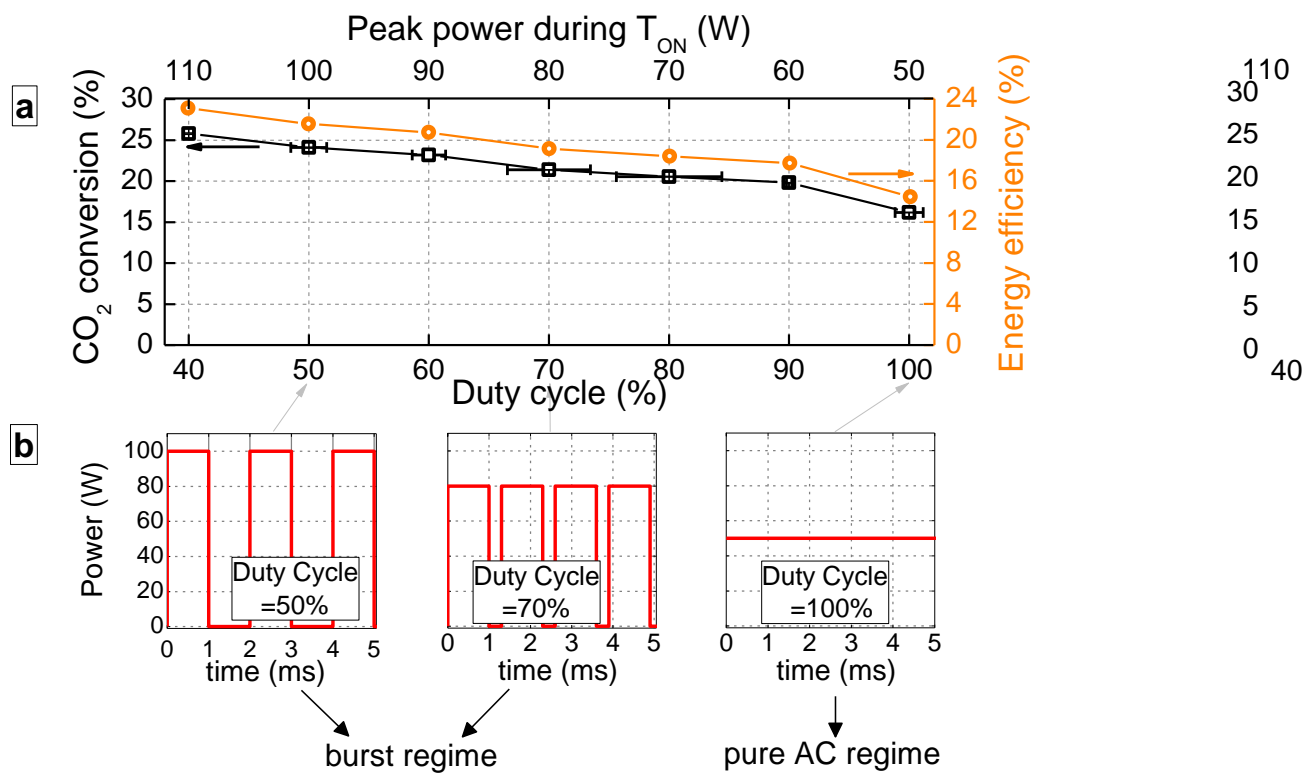


Figure 4 : (a) CO₂ conversion and energy efficiency as a function of the duty cycle – $P_{\text{applied}}=50 \text{ W}$, $f_{\text{signal}}=28.6 \text{ kHz}$, $f_{\text{repetition}}=400 - 900 \text{ Hz}$ and $\Phi(\text{CO}_2)=200 \text{ mL}_n\cdot\text{min}^{-1}$; (b) Power versus time for $D_{\text{cycle}}=50, 70$ and 100% .

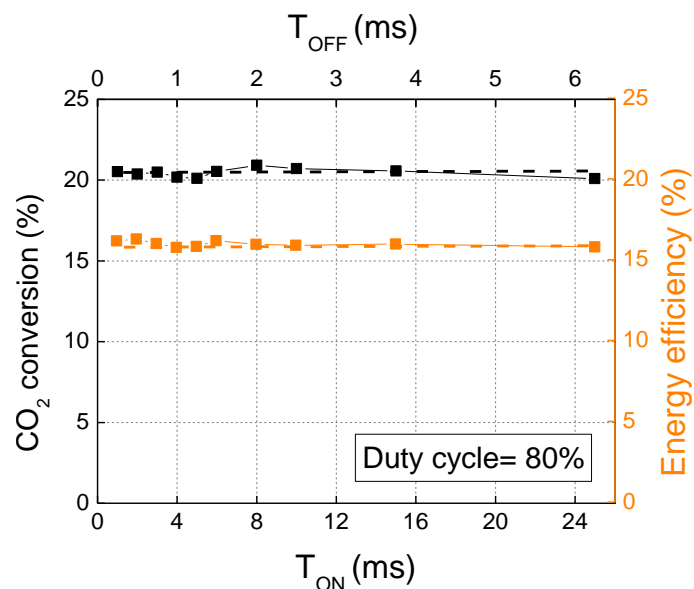


Figure 5 : CO₂ conversion and energy efficiency for a fixed duty cycle of 80%, as a function of T_{ON}. $P_{\text{applied}}=50 \text{ W}$, $f_{\text{signal}}=28.6 \text{ kHz}$ and $\Phi(\text{CO}_2)=200 \text{ mL}_n\cdot\text{min}^{-1}$.

Influence of the duty cycle on the discharge time-stability

Figure 6 shows the applied voltage as a function of the process time for several different duty cycles. The lower the duty cycle, the more stable and the higher is the voltage as a function of the process time. For $D_{\text{cycle}} < 70\%$, the voltage remains quite stable over time, e.g. at $D_{\text{cycle}} = 50\%$ it is close to 4400 V for all process times investigated (with only a small variation within the first 60 s). Therefore, the discharge appears as very stable in time. However, this time-stability gradually vanishes upon increasing D_{cycle} and in the pure AC mode ($D_{\text{cycle}} = 100\%$) the time dependency is quite significant: a drop of approximately 500 V is reached in 90 s, from 4400 V to a plateau at 3900 V. This transient regime is clearly visible in pure AC mode and is due to the dielectric properties of the barrier: it behaves as a capacitor with a dielectric relaxation time that is much longer than the period associated with the applied voltage. The dielectric relaxation time of the barrier t_{relax} is defined as the product of its dielectric permittivity ($\epsilon = \epsilon_0 \cdot \epsilon_r$) with its electrical resistivity (ρ), i.e., $t_{\text{relax}} = \epsilon_0 \cdot \epsilon_r \cdot \rho$, where $\epsilon_0 = 8.854 \cdot 10^{-12} \text{ F} \cdot \text{m}^{-1}$ and, in the case of alumina, $\epsilon_r = 9$ and $\rho \approx 10^{14} \Omega^{-1} \cdot \text{cm}^{-1}$ [68]. A rough estimation leads to $t_{\text{relax}} \approx 0.8 \text{ s}$. This value is much higher than the period of the applied voltage, which is $35.7 \mu\text{s}$ (since $f_{\text{signal}} = 28.6 \text{ kHz}$). In the first half period, the electrical charge deposited on the barrier cannot be entirely evacuated and partially remains in the second half period. This charge accumulation is negligible at the very beginning of the discharge ignition, but becomes significant in less than 30 seconds of operation (see Figure 6 for $D_{\text{cycle}} = 100\%$). Once the DBD is ignited, the barrier potential gradually increases, hence inducing a gradual decrease in V_{DBD} observed in Figure 6. This phenomenon is clearly less pronounced in the burst mode, where the T_{ON} times are too short to induce significant charge accumulations on the barrier. Moreover, the T_{OFF} times are much longer so that the sustaining voltage can remain closer to the ignition voltage [69]. The applied voltage (V_{DBD}) is expressed as the sum of two voltages, $V_{\text{DBD}} = V_{\text{pl,eff}} + V_{\text{diel}}$, where $V_{\text{pl,eff}}$ is the effective plasma voltage (in the gap) and V_{diel} is the voltage through the dielectric barrier. In a previous work [61], we have demonstrated that varying V_{DBD} for a fixed power induces variations of $V_{\text{pl,eff}}$ while V_{diel} remains almost unchanged. Here, decreasing the duty cycle at $P_{\text{applied}} = 50 \text{ W}$ only induces an increase in V_{DBD} , hence of $V_{\text{pl,eff}}$. As the gap is fixed, we can thus conclude that a stronger electric field is created as the plasma voltage is higher, and therefore there will be more energetic electrons. As a result, the burst mode can enhance the rates of electron impact excitation, ionization and dissociation of CO_2 and thus improve the conversion and energy efficiency of the CO_2 splitting process, as illustrated in Figure 4a. Hence, besides the higher peak power (due to the shorter duty cycle), this can also explain the higher CO_2 conversion in the burst mode.

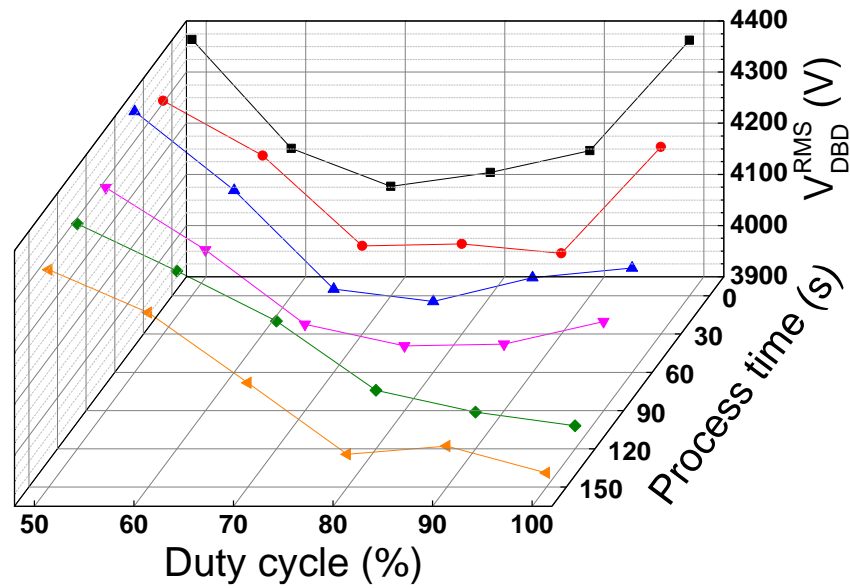


Figure 6 : RMS value of the DBD voltage as a function of the duty cycle and the process time. $P_{\text{applied}}=50 \text{ W}$, $f_{\text{signal}}=28.6 \text{ kHz}$, $f_{\text{repetition}}=500 - 900 \text{ Hz}$ and $\Phi(\text{CO}_2)=200 \text{ mL}_n\cdot\text{min}^{-1}$.

Influence of the duty cycle on the gas temperature

The influence of the duty cycle on the gas temperature is obtained from OES measurements. Ro-vibrational spectra of the Angstrom CO band are investigated, as well as the first positive system (FPS) of N_2 . In Figure 7, a linear decrease in gas temperature is observed when reducing the duty cycle, i.e. when increasing the T_{OFF} times. This is logical because in the burst mode, the gas can cool down. In the pure AC mode, where the CO_2 conversion is the lowest (16.1%), the gas temperature is the most elevated, suggesting that in general, a larger fraction of the power is used for gas heating, while a smaller fraction of the power leads effectively to CO_2 splitting. It should be noted that the gas temperature as obtained from the FPS of N_2 is about 100 K higher than the gas temperature obtained from the Angstrom CO band, over the entire range of duty cycles investigated. At this point, it is rather difficult to explain this difference since no other independent method (e.g. Doppler broadening, thermocouple, etc.) was used for the gas temperature determination in the plasma phase. Besides this, it is also worth mentioning that the line-ratio and synthetic spectra calculation methods are prone to errors as they assume a Boltzmann distribution and the validity of this assumption cannot be directly checked. Nevertheless, despite the difference of about 100 K, the trend of decreasing gas temperature with decreasing duty cycle is obvious from both curves, as shown in Figure 7.

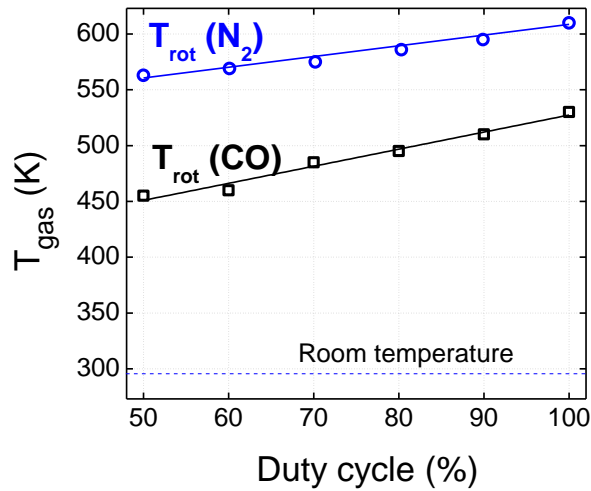


Figure 7 : Gas temperature as a function of duty cycle, as obtained from the FPS of N_2 and from the Angstrom CO band; $P_{applied}=50$ W, $f_{signal}=28.6$ kHz, $f_{repetition}=500 - 900$ Hz and $\Phi(CO_2)=200$ mL $_n$.min $^{-1}$.

Influence of the duty cycle on the electrical characteristics

Figure 8a shows a picture of the filaments in the DBD for two different duty cycles (50 and 100%). For this purpose, the alumina dielectric barrier is replaced by (transparent) borosilicate glass in order to visually observe these filaments. The pictures reveal an apparent larger number of filaments at $D_{cycle}=50\%$ than in pure AC mode ($D_{cycle}=100\%$). To study this in more detail, these pictures are correlated with their corresponding current oscillograms in Figure 8b, illustrating the currents peaks ($I_{DBD}=I_{pl}+I_{diel}$) during one AC period, i.e. 35.7 μ s. Note that the oscillogram corresponding to $D_{cycle}=50\%$ is a current profile related to $P_{plasma}=100$ W, as these oscillograms are recorded only during T_{ON} . A visual observation of these oscillograms indicates that the number of microdischarge pulses, as well as their lifetime and the current amplitude all increase from $D_{cycle}=100\%$ to $D_{cycle}=50\%$. Based on a numerical method explained in [61], we can also obtain more quantitative information on these microdischarges, such as their number (N_{md}), lifetime (L_{md}), charge (Q_{pl}) and conduction current (i_{pl}), from these current peaks.

These parameters are plotted as a function of duty cycle in Figure 9, as obtained for one AC period of 35.7 μ s. These results refer to the same applied power, and thus the peak power is higher in the burst mode. All these parameters are given in terms of (i) the same residence time (by taking into account T_{OFF} and thus, the electrical parameters are multiplied by D_{cycle}) and (ii) only the plasma ignition (without taking into account T_{OFF}). As observed in the figures, all of them are clearly changed in the burst mode and upon decreasing duty cycle. For instance in terms of same residence time, a reduction of D_{cycle} from 100 to 50% induces a decrease in the N_{md} from 390 to 340 per period. On the other hand, their mean lifetime (L_{md}) increases from 12.9 to 14.8 ns during the plasma ignition. A drastic rise of the plasma current i_{pl} (from 9 to 14 mA) is also observed when the duty cycle drops from 100 to 50%, which will result in a significant increase in electron density. The latter can be deduced from the rise in

plasma charge Q_{pl} in terms of a given residence time (from 0.3 to 0.5 μC). As the oscillograms are recorded during T_{ON} and the peak power becomes higher in the burst mode, the values of N_{md} , L_{md} , Q_{pl} and i_{pl} will increase upon increasing plasma power P_{plasma} (cf. our previous study on the effect of power on the electrical characteristics [61]). Therefore in the last paragraph, we will also compare the electrical characteristics in burst mode and pure AC mode, at the same delivered power during T_{ON} .

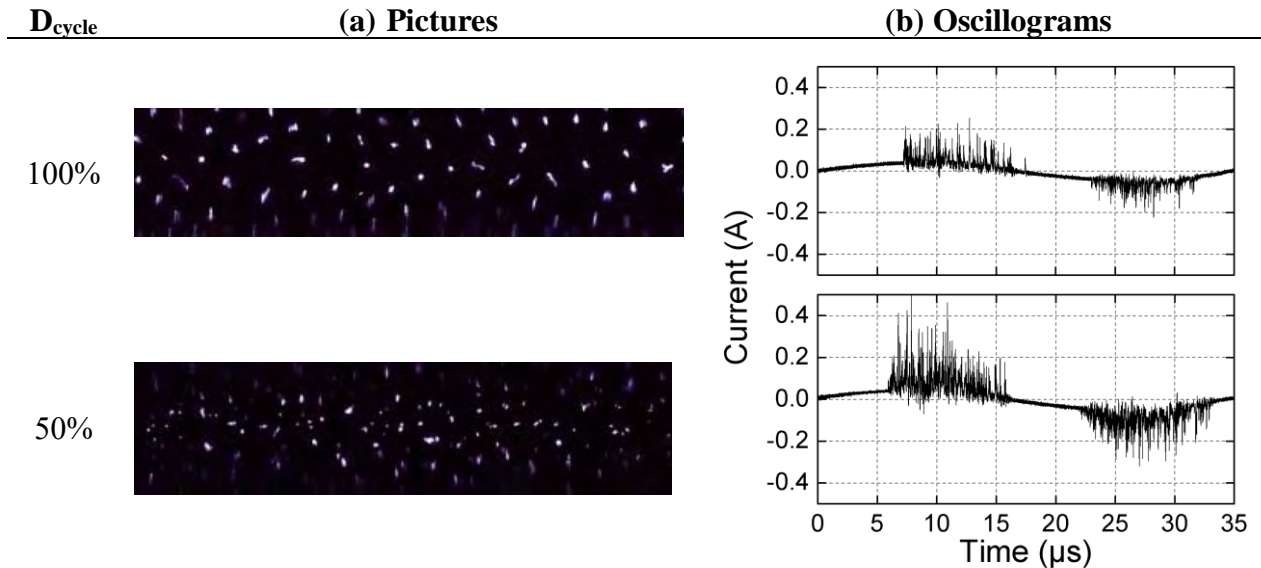


Figure 8 : (a) Pictures of the distribution of the microdischarges in the DBD, and (b) corresponding current oscillograms, for two different duty cycles (50 and 100%). $P_{applied}=50$ W, $f_{signal}=28.6$ kHz and $\Phi(\text{CO}_2)=200$ mL $_n$.min $^{-1}$. Camera aperture is 1/100 s.

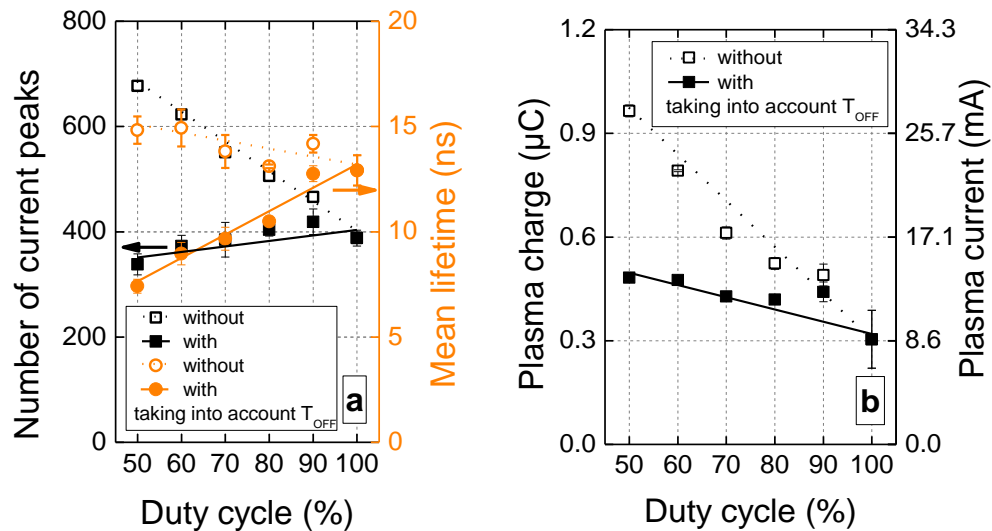


Figure 9 : (a) Number of microdischarge over one period (left axis) and their mean lifetime (right axis) versus duty cycle, and (b) plasma charge accumulation over one period (left axis) and plasma current (right axis) versus duty cycle. All electrical parameters are given either considering a total residence time (solid lines) or T_{ON} (dot lines). $P_{applied}=50$ W, $f_{signal}=28.6$ kHz, $f_{repetition}=500 - 900$ Hz and $\Phi(\text{CO}_2)=200$ mL $_n$.min $^{-1}$.

Correlation between microdischarge parameters and energy efficiency of burst and pure AC mode

Figure 10 depicts the number of microdischarges, their mean lifetime and the plasma charge and current for pure AC (values adopted from our previous paper [61]) and burst mode, as a function of plasma power. Several observations are noteworthy:

- (i) In pure AC mode (i.e., all corresponding to $D_{\text{cycle}}=100\%$), the plasma power corresponds to the applied power ($P_{\text{plasma}}=P_{\text{applied}}$). Thus, the plasma power is increased by tuning the applied power.
- (ii) The plasma power in burst mode corresponds to $P_{\text{plasma}}=P_{\text{applied}}/D_{\text{cycle}}$ where P_{applied} is fixed (50 W). The plasma power (delivered during T_{ON}) is increased by tuning the duty cycle. For example, $P_{\text{plasma}}=50$ W at $D_{\text{cycle}}=100\%$ and ca. 100 W at $D_{\text{cycle}}=50\%$. The corresponding duty cycles are indicated next to the black points in the figure.
- (iii) All the aforementioned results in the article can only be compared to the black curves in Figure 10, i.e. for $P_{\text{applied}} = 50\text{W}$.

In Figure 10a, the number of microdischarges (N_{md}) is determined for $T_{\text{ON}}=1\text{ms}$ and multiplied by the duty cycle to always consider the same residence. Whatever the plasma power, N_{md} is higher in pure AC mode than in burst mode. For instance, at $P_{\text{plasma}}=100$ W, the burst mode (for 50% duty cycle) allows the formation of 340 microdischarges per period at $P_{\text{applied}}=50$ W, while 480 microdischarges are obtained at $P_{\text{applied}}=100\text{W}$ in the pure AC mode. The burst mode presents a lower number of microdischarges, but the applied power is twice as low, which is thus more interesting in terms of energy efficiency. For the same plasma power, a lower N_{md} means a higher plasma voltage ($V_{\text{pl,eff}}$), hence a higher electric field and thus higher electron temperatures. The filaments are generated in a higher electric field, leading to a higher efficiency of the CO_2 conversion.

Figure 10b and Figure 10c give information on the microdischarge lifetime as well as the electrical charge and current versus plasma power. These electrical parameters obviously increase with plasma power and the differences between pure AC mode and burst mode are noteworthy: Q_{pl} is indeed lowered in the burst mode. In terms of plasma ignition, the two modes result in a quite similar plasma charge accumulation and plasma current, when operating at the same delivered power, as observed in Figure 10c. As the plasma charge is directly related to the plasma current, the electron densities might be very similar in both modes during T_{ON} .

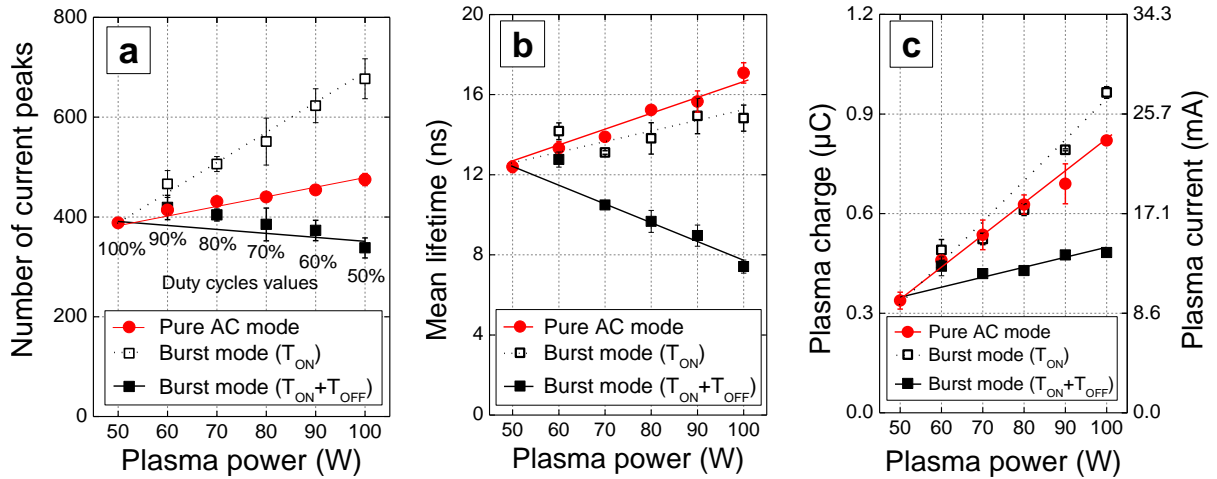


Figure 10 : (a) Number of microdischarges during one period, (b) microdischarges mean lifetime, and (c) plasma charge accumulation/plasma current during one period, as a function of the plasma power. P_{applied} is fixed in the burst mode while D_{cycle} is varied. P_{applied} is changed in the pure AC mode (electrical parameter values are adopted from our previous article [61]). $f_{\text{signal}}=28.6 \text{ kHz}$ and $\Phi(\text{CO}_2)=200 \text{ mL}_n \cdot \text{min}^{-1}$.

Conclusion

We studied the CO_2 dissociation in a cylindrical DBD reactor operating at atmospheric pressure, focusing on the comparison between pure AC mode and so-called burst mode, in which an AC voltage is only applied during a certain amount of time. The AC voltage signal has a period of $35.7 \mu\text{s}$ (corresponding to an applied frequency of 28.6 kHz) while the time width of the voltage burst (i.e., T_{ON}) is 1 ms and the T_{OFF} period varies between 0 and 1.5 ms (corresponding to duty cycles (D_{cycle}) between 100 and 40%, respectively). Note that $D_{\text{cycle}}=100\%$ corresponds to the AC mode. Comparing this usual AC mode with the burst mode leads us to conclude that the CO_2 conversion is significantly higher in the burst mode, and it increases from 16.1 to 25.8% upon decreasing D_{cycle} from 100 to 40%. Likewise, the corresponding energy efficiency of the CO_2 splitting rises from 14.5 to 23.1%.

To explain the impact of the plasma electrical properties on the CO_2 conversion, a detailed electrical characterization is carried out in the burst mode for various duty cycles. A comparison between the burst and pure AC mode at the same plasma powers reveals that the plasma currents and microdischarge mean lifetimes are quite similar, while the number of microdischarges appears lower in the burst mode.

We have also investigated the time-variation of the applied DBD voltage in the two modes: it is much more stable and constant as a function of processing time at low duty cycle (burst mode), whereas the voltage always drops at duty cycles higher than 70%, or in the pure AC mode. Also, we have shown that for a same plasma power, a lower N_{md} means a higher plasma voltage ($V_{\text{pl,eff}}$), hence a higher electric field and thus higher electron temperatures. The filaments are generated in a higher electric field, leading to a higher efficiency of the CO_2 conversion.

Finally, OES measurements have shown that the burst mode leads to a lower gas temperature, indicating that a smaller fraction of the power is used for gas heating, so that a larger fraction can be dedicated to the CO₂ dissociation.

Overall we can conclude that the so-called burst mode yields a significantly higher CO₂ conversion for the same applied power as in a usual AC mode, which is very promising in terms of applications.

Acknowledgements

The authors acknowledge financial support from the IAP-VII/12, P7/34 (Inter-university Attraction Pole) program ‘PSI-Physical Chemistry of Plasma-Surface Interactions’, financially supported by the Belgian Federal Office for Science Policy (BELSPO). A. Ozkan would also like to thank financial support given by ‘Fonds David et Alice Van Buuren’.

Bibliography

- [1] Pachauri R K, Allen M, Barros V, Broome J, Cramer W, Christ R, Church J, Clarke L, Dahe Q and Dasgupta P 2014
- [2] Feely R A, Doney S C and Cooley S R 2009 *Oceanography* **22** (4) 36-47
- [3] Bindoff N L, Willebrand J, Artale V, Cazenave A, Gregory J M, Gulev S, Hanawa K, Le Quéré C, Levitus S, Nojiri Y, Shum C, Talley L and Unnikrishnan A 2007 *Observations: oceanic climate change and sea level* (Cambridge University Press)
- [4] Rahmstorf S 2007 *Science* **315** (5810) 368-370
- [5] Jones P, Trenberth K, Ambenje P, Bojariu R, Easterling D, Klein T, Parker D, Renwick J, Rusticucci M and Soden B 2007 *In Climate Change 2007: The Physical Science Basis Contribution of Working Group I to the Fourth Assessment Report of the Intergovernmental Panel on Climate Change* 235-336
- [6] Grinsted A, Moore J C and Jevrejeva S 2010 *Climate Dynamics* **34** (4) 461-472
- [7] Church J A and White N J 2006 *Geophysical research letters* **33** (1)
- [8] Gibbins J and Chalmers H 2008 *Energy Policy* **36** (12) 4317-4322
- [9] Thomas D C and Benson S M 2015 *Carbon Dioxide Capture for Storage in Deep Geologic Formations-Results from the CO₂ Capture Project: Vol 2-Geologic Storage of Carbon Dioxide with Monitoring and Verification* (Elsevier)
- [10] Haszeldine R S 2009 *Science* **325** (5948) 1647-1652
- [11] Aresta M 2010 *Carbon dioxide as chemical feedstock* (John Wiley & Sons)
- [12] Christophe J, Doneux T and Buess-Herman C 2012 *Electrocatalysis* **3** (2) 139-146
- [13] Fridman A 2008 *Plasma chemistry* (Cambridge university press)
- [14] Liu C-j, Xu G-h and Wang T 1999 *Fuel Processing Technology* **58** (2) 119-134
- [15] Zou J-J, Zhang Y-p, Liu C-J, Li Y and Eliasson B 2003 *Plasma Chemistry and Plasma Processing* **23** (1) 69-82
- [16] Bo Z, Yan J, Li X, Chi Y and Cen K 2008 *International Journal of Hydrogen Energy* **33** (20) 5545-5553
- [17] Zhang Y-p, Li Y, Wang Y, Liu C-j and Eliasson B 2003 *Fuel Processing Technology* **83** (1) 101-109
- [18] Tao X, Bai M, Li X, Long H, Shang S, Yin Y and Dai X 2011 *Progress in Energy and Combustion Science* **37** (2) 113-124

- [19] Goujard V, Tatibouët J-M and Batiot-Dupeyrat C 2009 *Applied Catalysis A: General* **353** (2) 228-235
- [20] Li Y, Liu C-J, Eliasson B and Wang Y 2002 *Energy & fuels* **16** (4) 864-870
- [21] Ozkan A, Dufour T, Arnoult G, De Keyzer P, Bogaerts A and Reniers F 2015 *Journal of CO2 utilization* **9** 74-81
- [22] Paulussen S, Verheyde B, Tu X, De Bie C, Martens T, Petrovic D, Bogaerts A and Sels B 2010 *Plasma Sources Science and Technology* **19** (3) 034015
- [23] Mei D, He Y L, Liu S, Yan J and Tu X 2015 *Plasma Processes and Polymers*
- [24] Brehmer F, Welzel S, van de Sanden M and Engeln R 2014 *Journal of Applied Physics* **116** (12) 123303
- [25] Jiang H, Shao T, Zhang C, Niu Z, Yu Y, Yan P and Zhou Y 2011 *Plasma Science, IEEE Transactions on* **39** (11) 2076-2077
- [26] Williamson J M, Trump D D, Bletzing P and Ganguly B N 2006 *Journal of Physics D: Applied Physics* **39** (20) 4400
- [27] Shao T, Yu Y, Zhang C, Zhang D, Niu Z, Wang J, Yan P and Zhou Y 2010 *Dielectrics and Electrical Insulation, IEEE Transactions on* **17** (6) 1830-1837
- [28] Zhang C, Shao T, Ma H, Ren C, Yan P and Zhou Y 2014 *Plasma Science, IEEE Transactions on* **42** (10) 2354-2355
- [29] Shao T, Zhang C, Zhou Y, Xie Q, Zhou Z and Yan P 2014 *Plasma Science, IEEE Transactions on* **42** (10) 2408-2409
- [30] Zhang C, Shao T, Wang R, Zhou Z, Zhou Y and Yan P 2014 *Physics of Plasmas (1994-present)* **21** (10) 103505
- [31] Jiang T, Li Y, Liu C-j, Xu G-h, Eliasson B and Xue B 2002 *Catalysis Today* **72** (3) 229-235
- [32] Ghorbanzadeh A, Lotfalipour R and Rezaei S 2009 *international journal of hydrogen energy* **34** (1) 293-298
- [33] Ghorbanzadeh A and Modarresi H 2007 *Journal of applied physics* **101** (12)
- [34] Yao S, Okumoto M, Nakayama A and Suzuki E 2001 *Energy & fuels* **15** (5) 1295-1299
- [35] Shao T, Zhang C, Fang Z, Yu Y, Zhang D, Yan P, Zhou Y and Schamiloglu E 2013 *Plasma Science, IEEE Transactions on* **41** (10) 3069-3078
- [36] Jiang Z and Jiang Z-j 2012 *International Journal of Hydrogen Energy* **37** (15) 11276-11289
- [37] Merche D, Vandencastele N and Reniers F 2012 *Thin Solid Films* **520** (13) 4219-4236
- [38] Walsh J L and Kong M G 2007
- [39] Ndong A A a, Zouzou N, Benard N and Moreau E 2013 *Journal of Electrostatics* **71** (3) 246-253
- [40] Shao T, Yang W, Zhang C, Fang Z, Zhou Y and Schamiloglu E 2014 *EPL (Europhysics Letters)* **107** (6) 65004
- [41] Song H K, Lee H, Choi J-W and Na B-k 2004 *Plasma chemistry and plasma processing* **24** (1) 57-72
- [42] Lee H, Lee C-H, Choi J-W and Song H K 2007 *Energy & fuels* **21** (1) 23-29
- [43] Khacef A and Cormier J M 2006 *Journal of Physics D: Applied Physics* **39** (6) 1078
- [44] Ghorbanzadeh A, Norouzi S and Mohammadi T 2005 *Journal of Physics D: Applied Physics* **38** (20) 3804
- [45] Benard N and Moreau E 2010 *Journal of Physics D: Applied Physics* **43** (14) 145201
- [46] Gejima H, Takinami R, Fukagata K, Mitsumoji T, Sueki T and Ikeda M 2015 *Journal of Fluid Science and Technology* **10** (1) JFST0006-JFST0006
- [47] Benard N, Sujar-Garrido P, Bayoda K D, Bonnet J-P and Moreau E, editors. Pulsed dielectric barrier discharge for manipulation of turbulent flow downstream a backward-

- facing-step. 52nd AIAA Aerospace Sciences Meeting, National Harbor, MD, USA, January; 2014.
- [48] Benard N, Bonnet J-P, Touchard G and Moreau E 2008 *AIAA journal* **46** (9) 2293-2305
- [49] Roupasov D, Nikipelov A, Nudnova M and Starikovskii A Y 2009 *AIAA journal* **47** (1) 168-185
- [50] Little J, Nishihara M, Adamovich I and Samimy M 2010 *Experiments in fluids* **48** (3) 521-537
- [51] Benard N and Moreau E 2014 *Experiments in Fluids* **55** (11) 1-43
- [52] Choi J-h, Huh D-y and Kim Y-s, editors. The improved burst mode in the stand-by operation of power supply. Applied Power Electronics Conference and Exposition, 2004 APEC'04 Nineteenth Annual IEEE; 2004: IEEE.
- [53] Andersen T, Rødgaard M S, Andersen M A, Thomsen O C, Lorenzen K, Mangeot C and Steenstrup A, editors. Integrated high voltage power supply utilizing burst mode control and its performance impact on dielectric electro active polymer actuators. 13th International Conference on New Actuators and 7th International Exhibition on Smart Actuators and Drive Systems; 2012.
- [54] Jiang H, Shao T, Zhang C, Li W, Yan P and Schamiloglu E, editors. Study on QV Lissajous figures in nanosecond-pulsed surface discharge. Power Modulator and High Voltage Conference (IPMHVC), 2012 IEEE International; 2012: IEEE.
- [55] Jiang H, Shao T, Zhang C, Li W, Yan P, Che X and Schamiloglu E 2013 *Dielectrics and Electrical Insulation, IEEE Transactions on* **20** (4) 1101-1111
- [56] Shao T, Jiang H, Zhang C, Yan P, Lomaev M I and Tarasenko V F 2013 *EPL (Europhysics Letters)* **101** (4) 45002
- [57] Manley T 1943 *Transactions of the electrochemical society* **84** (1) 83-96
- [58] Wagner H-E, Brandenburg R, Kozlov K, Sonnenfeld A, Michel P and Behnke J 2003 *Vacuum* **71** (3) 417-436
- [59] Aerts R, Martens T and Bogaerts A 2012 *The Journal of Physical Chemistry C* **116** (44) 23257-23273
- [60] Aerts R, Somers W and Bogaerts A 2015 *ChemSusChem* **8** (4) 702-716
- [61] Ozkan A, Dufour T, Silva T, Britun N, Snyders R, Bogaerts A and Reniers F 2016 *Plasma Sources Science and Technology* **25** (2) 025013
- [62] Ozkan A, Dufour T, Bogaerts A and Reniers F 2016 *submitted to Plasma Sources Science and Technology*
- [63] Masoud N, Martus K, Figus M and Becker K 2005 *Contributions to Plasma Physics* **45** (1) 32-39
- [64] Silva T, Britun N, Godfroid T and Snyders R 2014 *Plasma Sources Science and Technology* **23** (2) 025009
- [65] Reginald William Blake P and Alfred Gordon G 1976 *The identification of molecular spectra*. 4th edition ed. London/New York (Chapman and Hall/Wiley)
- [66] Britun N, Godfroid T, Konstantinidis S and Snyders R 2011 *Applied Physics Letters* **98** (14) 141502
- [67] Britun N, Godfroid T and Snyders R 2012 *Plasma Sources Science and Technology* **21** (3) 035007
- [68] Manceau J-P. Etude du phénomène de relaxation diélectrique dans les capacités Métal-Isolant-Métal: Université Joseph-Fourier-Grenoble I; 2008.
- [69] Valdivia-Barrientos R, Pacheco-Sotelo J, Pacheco-Pacheco M, Benitez-Read J and López-Callejas R 2006 *Plasma sources science and technology* **15** (2) 237