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Characteristics of a novel nanosecond DBD microplasma reactor for flow applications

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Abstract

We present a novel microplasma flow reactor using a dielectric barrier discharge (DBD) driven by repetitive nanosecond high-voltage pulses. Our DBD-based geometry can generate a non-thermal plasma discharge at atmospheric pressure and below in a regular pattern of micro-channels. This reactor can work continuously up to about 100 min in air, depending on the pulse repetition rate and operating pressure. We here present the geometry and main characteristics of the reactor. Pulse energies of 1.46 and 1.3 $\mu$J per channel at atmospheric pressure and 50 mbar, respectively, have been determined by time-resolved measurements of current and voltage. Time-resolved optical emission spectroscopy measurements have been performed to calculate the relative species concentrations and temperatures (vibrational and rotational) of the discharge. The effects of the operating pressure and flow velocity on the discharge intensity have been investigated. In addition, the effective reduced electric field strength $(E/N)_{\text{eff}}$ has been obtained from the intensity ratio of vibronic emission bands of molecular nitrogen at different operating pressures and different locations. The derived $(E/N)_{\text{eff}}$ increases gradually from about 550 to 4600 Td when decreasing the pressure from 1 bar to 100 mbar. Below 100 mbar, further pressure reduction results in a significant increase in $(E/N)_{\text{eff}}$ up to about 10000 Td at 50 mbar.

Keywords: microplasma, DBD microplasma, atmospheric pressure plasma, plasma spectroscopy

1. Introduction

Plasma flow reactors can be used to chemically process a gas. They are of interest because of their high energy efficiency and high selectivity. Therefore, they have a strong potential in applications such as surface treatment [1], thin-film deposition [2], detoxification of gaseous pollution [3, 4], plasma medicine [5] and ozone generation [6]. Many different designs have been tried to generate a stable non-thermal plasma discharge in a flow reactor at pressures up to one atmosphere. Some of these designs are plasma jets [7, 8], packed-bed plasma reactors [9, 10], pulsed-corona plasma reactors [11–13] and atmospheric pressure glow discharges [14].

One of the most extensively studied configurations is an array of so-called microhollow cathode discharges (MHCDs), which has the ability to produce a high-density plasma discharge in a relatively small volume (characteristic dimension of about $10^{-4}$ m) [15–17]. Based on that, compared to other types of flow reactors, MHCD ensures a maximum interaction between the plasma discharge and the processed gas. However, two conditions should be fulfilled to maintain a discharge in MHCD devices. First, the applied voltage should exceed the breakdown voltage and be able to ignite the discharge for the given gas and pressure [18]. Second, according to the Allis–White similarity law, the product of the pressure $(p)$ and the aperture diameter of the cathode $(D)$ should fall in the typical operating range for MHCD, which is $0.1–13$ mbar-cm depending on the gas, electrodes and geometry [19]. These conditions pose some constraints regarding scaling-up the diameter of the MHCD devices at high pressures. Based on the upper limit of the product $(p \cdot D)$, atmospheric pressure operation in air would lead to a maximum diameter $(D)$ of 100 $\mu$m. However, in many applications, especially for gas...
treatment, it is desirable to increase the diameter to reduce the pressure drop for the same flow condition.

One of the most successful ways to increase the hole diameter of the reactor is by using a dielectric barrier discharge (DBD) \([20, 21]\) driven by nanosecond high-voltage pulses. The surface discharge deposited by the (pulsed) discharge counteracts the applied field, thereby self-limiting the discharge. Together with the nanosecond pulses, this helps to generate a uniform discharge distribution and reduces the chance of glow-to-arc transition. In addition, using nanosecond repetitive high-voltage pulses provides highly energetic electrons and more chemically excited species compared to AC power sources due to the rapid ionization process, with a power consumption that is about one order of magnitude lower \([22, 23]\). The increase of the electron energy comes without a considerable increase of the gas temperature, thereby reducing undesired thermal effects of the plasma discharge on the treated gas.

In this paper, we introduce a new geometry, which utilizes the DBD method to sustain a non-thermal plasma discharge in an array of 400 \(\mu\)m diameter channels perforating a 1.5 mm thick dielectric slab with embedded electrodes. The discharge is powered by nanosecond high-voltage pulses and is operated at pressures up to one atmosphere, which corresponds to a pressure \((p)\) times diameter \((D)\) of 40 mbar-cm. The main development in this study is that the high-voltage electrode is fully embedded within the dielectric substrate. This allows us to generate a non-thermal plasma discharge at atmospheric pressure in channels larger than the maximum possible diameter for MHCDs at the same pressure.

The geometry consists of 363 channels of 400 \(\mu\)m in diameter placed in parallel. Each channel has two sets of electrodes, stacked to work (optionally) in series to increase the discharge volume in order to increase the efficiency of the gas-discharge interaction. Figure 1 shows a top-view luminosity image of a plasma discharge in this DBD microplasma reactor, with 3.4 l min\(^{-1}\) air flow upwards, out of the plane of the paper, at atmospheric pressure, 5 kV voltage pulse with a repetition rate of 3 kHz, recorded by a digital DSLR camera (Nikon D5100).

The non-thermal nature and the high electron energy of the produced discharge make this geometry very promising for applications such as plasma-assisted combustion, chemical treatment, light sources, plasma medicine, pollutant control and many others.

In this paper, we present the geometry, material and fabrication process of the DBD microplasma reactor as well as the characteristics of the nanosecond pulsed plasma produced by it. Time-resolved optical spectroscopy is used to investigate the discharge emission and its temperatures. ICCD images are analyzed to identify the discharge structures at different operating pressures. The relation between the effective reduced electric field strength \((E/N)_{\text{eff}}\) and operating pressure and the effect of flow velocity on discharge emission are also presented. Finally, the reactor lifetime is determined for different operating pressures and pulse repetition rates.

An example of the use of our reactor is given in figure 2 where it stabilizes a methane-air flame in order for us to study the effect of the plasma discharge on combustion characteristics. The preliminary results of this study show a significant increase, reaching up to 100% in some cases, in the laminar burning velocity as a result of the plasma discharge. Results of this combustion study will be published within a different framework.

2. Experimental setup

2.1. Reactor geometry, materials and fabrication

As shown in figure 3, the new plasma device is a 1.5 mm thick dielectric slab perforated by a hexagonal array of 363 channels. The channels have a diameter of 0.4 mm and a pitch of 1.0 mm. The outer diameter of the reactor is 70 mm, while the effective diameter of the flow area is 20 mm. The reactor geometry includes four layers of copper electrodes (A, B, C and D) in figure 3B. The high-voltage electrodes (B and C) are embedded into the dielectric material. The grounded electrodes (A and D) are located on the top and bottom of the reactor.
surfaces of the reactor. The outer and the inner copper layers have a diameter of 40 and 30 mm, respectively. All electrodes are perforated in the same hexagonal pattern as the dielectric, but the holes are twice as large as those in the dielectric. The device can work in two modes: single-layer discharge mode and double-layer discharge mode. In the single-layer discharge mode, only electrodes A and B are activated (see figure 5 for the equivalent circuit diagram), while electrodes C and D are kept floating. All the work presented in this study has been done in the single-layer discharge mode.

The cathode and anode are separated by a dielectric layer with a thickness of 0.36 mm. The outer (grounded) and inner (high-voltage) electrodes are made of copper with thicknesses of 18 and 38 μm, respectively. The separation distance between the two embedded electrodes is 0.6 mm, while the total thickness of the reactor is 1.5 mm. The width and thickness of the embedded electrodes have been calculated to safely withstand the peak current of the discharge pulses, which is 205 mA per channel for a duration of 10 ns, as shown in figure 7(a).

We came to this design by performing a geometrical parameter study to (i) determine the minimum dielectric thickness in order to maximize the discharge strength and (ii) guide most of the discharge to the inside of the holes instead of the top or bottom reactor surfaces, to get an efficient interaction with the flowing gas. From this study we found that the dielectric thickness between the anode and the wall of the hole, \( L_1 \) in view (B) of figure 3, should be less than the axial dielectric thickness between the anode and cathode \( L_2 \).

The dielectric material of the reactor is a composite of woven electrical grade fiberglass and epoxy resin. It has a dielectric constant \( D_r \) of 4.17 ± 0.05 (at 1 GHz/23 °C), a thermal conductivity of 0.4 W m K \(^{-1}\), a dielectric breakdown strength of 32 kV mm \(^{-1}\), a thermal expansion coefficient of 14 ppm/°C and a glass transition temperature of 180 °C. This material has a stable performance over a wide range of frequencies and temperatures, as well as good machinability.

In order to reduce the chance of micro-filamentary discharges on the reactor surface, which would reduce the reactor lifetime as a result of excessive heat, the outer copper electrodes have been coated with a nickel/immersion gold layer, which gives a surface roughness of 0.5 μm. In some applications, which are accompanied by thermal radiation, such as plasma-assisted combustion or chemical vapor deposition, the copper layers also act as protective layers to conduct away excessive heat from the process. This prevents overheating of the dielectric material, which can lead to dielectric failure.

The plasma reactor is manufactured by a multi-step UV lithography process. First, the desired copper patterns are precisely defined on two dielectric slabs with copper layers pre-bonded onto each side. These correspond to the outer and inner electrode pairs and will be stacked on top of each other later in the process. Then, UV light is used to harden a photoresist layer on the desired copper tracks, after which an alkaline solution is sprayed over the panel to etch away the undesired copper. Next, the two panels are stacked together bonded by another epoxy resin layer and thermally pressed to guarantee a permanent bond without any air gap. Then, the copper tracks on the upper and lower layers are coated with a nickel/immersion gold layer. Finally, the channels are drilled with a laser drilling machine to ensure clean hole walls and a uniform pattern.

2.2. Flow setup

A schematic representation of the experimental setup is shown in figure 4. The setup consists of three main parts; the plenum chamber at the bottom, a water jacket in the middle and the microplasma reactor on top. The plenum chamber and the water jacket are made of steel and brass, respectively. The
The purpose of these parts is to create a uniform air flow towards the plasma reactor. The gas inlet port is located at the bottom of the plenum chamber. A perforated plate at 20 mm from the bottom works as a flow straightener. The holes in the perforated plate (1.0 mm in diameter and 2.0 mm in pitch) cover the entire area of the plate. In case of reduced pressure, a vacuum chamber of 20 mm height and 60 mm in diameter is mounted on top of the plasma reactor. The top flange of the chamber includes a fused quartz window with a thickness of 4 mm, through which the discharge emission can be viewed. The reactor temperature is controlled by means of the water jacket underneath the reactor. The water inlet and outlet are connected to a heater and thermostat to control the water temperature. This water circuit can heat up and control the upper surface of the reactor up to 80°C.

The air flow is controlled by a mass flow controller (MFC; Bronkhorst model F-202CV). A cylindrical buffer vessel is placed before the MFC to damp pressure fluctuations. In addition, an air filter is installed just before the MFC to avoid contamination. To ensure purity, synthetic air of grade 4.0 has been used. A vacuum pump model Busch R5 KB 0010, with a minimum pressure of 20 mbar and flow of 10 m³ h⁻¹, has been used to evacuate the vacuum chamber to the desired pressure.

2.3. Electrical characteristics

Figure 5 shows the equivalent electrical circuit diagram as used with single-layer operation (red box) and double-layer operation (blue box) for the DBD microplasma reactor. In the case of double-layer operation, the total voltage \( V_t \) is the same for the two layers, while the current is divided evenly between the two layers. When the voltage exceeds the breakdown voltage, indicated as an ignition switch in the figure, the current \( I_D \) will pass through the discharge gap to initiate the plasma discharge.

The high-voltage pulses are produced by a Mega-Impulse semiconductor-based pulse generator, model NPG-6/15 k. Positive polarity pulses of 4–7 kV in amplitude, 25 ns in duration and 10 ns rise time are created at a pulse repetition frequency in the range of 0–10 kHz. The pulse generator is...
externally triggered by a FeelTech frequency generator model FY2102S. Voltage and current are simultaneously monitored by a Lecroy waverunner 44MXi-A oscilloscope with an analog bandwidth of 400 MHz and sampling rate of 5 GS s⁻¹.

The voltage is recorded by a North Star high-voltage probe model PVM-4 with a sensitivity of 1 V/1 kV, a bandwidth of 120 MHz and 2 ns rise time. The current through the electrodes is measured by a Bergoz current transformer model CT-F0.5 with a 1.75 ns rise time and a sensitivity of 0.5 V/A voltage-to-current conversion located on the ground cable.

For precise calculation of the pulse energy, it is important to make sure that the voltage and current signals are synchronized. This was achieved by matching the calculated capacitive current from the voltage signal and the measured current for a pure capacitive case. To ensure a pure capacitive current, the applied voltage across the electrodes is set to the minimum possible value of the power supply, which is 4 kV with a pulse repetition rate of 10 Hz. Also, the capacitance between the electrodes has been increased by filling the channels and covering the surfaces of the reactor with silicon material, preventing a discharge from developing. The waveshape of the expected capacitive current ($I_{cap}$) was then calculated by taking the time-derivative of the voltage signal ($V$) multiplied by the modified capacitance of the device ($C_{mod}$):

$$I_{cap} = C_{mod} \frac{dV}{dt}$$

Figure 6 shows the measured voltage and current waveforms and the calculated capacitive current, for the modified reactor. A time delay between the measured current and the calculated capacitive current of 3 ns has been found. This time delay has been added to all the measured current signals to adjust for the phase-shift.

The reciprocal of the effective capacitance $C_{total}$ is the sum of the reciprocal of two components, the reactor capacitance $C_R$ and the cable capacitance $C_{cable}$. Both the reactor and cable capacitances have been measured by a Fluke RCL meter model PM6303, resulting in $C_R = 94$ pF and
Table 1. Blaze wavelength and equivalent FWHM of the slit function for a 20 μm wide slit of the gratings used in this experiment.

| Grating (grooves/mm) | 300 | 1800 | 3600 |
|----------------------|-----|------|------|
| Blaze wavelength (nm) | 300 | 200  | 300  |
| FWHM (nm)            | 0.58| 0.084| 0.037|

$C_{\text{cable}} = 200 \text{ pF}$. So, the equivalent total capacitance is $C_{\text{total}} = 64 \text{ pF}$.

2.4. Optical emission spectroscopy setup

Optical emission spectroscopy has been performed with a SpectraPro 2500i spectrometer with a focal distance of 0.5 m, fitted with three gratings of 300, 1800 and 3600 grooves/mm. Table 1 shows the blaze wavelengths and the equivalent full widths at half-maximum (FWHM) of the slit functions (approximately a Gaussian shape) for a 20 μm wide slit measured at 355 nm for these gratings. The emission spectra from the plasma discharge have been recorded using a 1024 × 1024 pixel ICCD camera (Princeton Instruments PIMax 3) on the exit port of the spectrogaph. The spectral line wavelength of the entire optical system is calibrated using the 253.65, 435.83 and 579.07 nm lines of a low-pressure pencil-type mercury lamp [24], model ORIEL 6035. A UV-Nikkor lens with 105 mm focal length and f/4.5 is attached to the spectrometer entrance slit to collect the emitted light from the plasma discharge.

Synchronization between the voltage pulse signal and the ICCD camera gate was achieved by a triggering signal from the high-voltage pulse generator, which was adjusted by the internal delay generator of the ICCD camera. A 200 ns time delay has been observed between the rising edge of the high-voltage pulse and the onset of the plasma discharge. This time delay comes from three sources: (1) the high-voltage pulse generator delay, (2) signal delay due to 3 m cable length and (3) the charging of the reactor capacitance. The digital images have been recorded using a commercial 16.2 mega-pixel digital camera Nikon D5100 fitted with a micro-Nikkor lens with a 60 mm focal length and f/2.8. The high-speed images have been recorded using a FASTCAM Mini UX100 camera fitted with an Invisible Vision intensifier model UVi 1850-05 and a 60 mm Nikkor micro lens. The synchronization between the image intensifier and the high-voltage pulses was achieved by a BNC-575 pulse-delay generator.

3. Results and discussion

3.1. Pulse characteristics and energy calculation

Figure 7(a) shows the current and voltage waveforms, after synchronization, of a single pulse for a single-layer operation in stagnant air at atmospheric pressure (solid lines) and at 50 mbar (dashed lines). The voltage signal shows a positive polarity incident pulse with a 5 kV peak followed by a negative polarity reflected pulse with a 1.75 kV peak. The rise time of the incident pulse is about 10 ns. The current signal shows the same behavior, with a positive peak of 75 A, corresponding to 205 mA per channel, followed by a negative peak of 58 A. The FWHM duration of the current signal is 7 ns and it has a rise time of 4.4 ns.

The instantaneous pulse power $P_{\text{pulse}}$ (blue lines in figure 7(b)) has been obtained by multiplying the measured voltage $V$ with the current $I$ (after adjusting the time delay). The power has been integrated over the pulse duration to get the total energy $E_{\text{pulse}}$ deposited in the plasma discharge:

$$E_{\text{pulse}} = \int_{t_1}^{t_2} I \cdot V dt.$$  

As shown in figure 7(b) (red lines), the total energy per pulse is about 0.53 mJ/pulse at atmospheric pressure and 0.47 mJ/pulse at 50 mbar. These values correspond to pulse energy of 1.46 and 1.3 μJ per channel, respectively. Also, it has been observed that the incident pulse is followed by a reflected pulse after about 40 ns, followed by some ripples. This behavior has been validated by time-resolved measurement of the discharge emission, reported in section 3.3.2 below.

3.2. Discharge structure

Figure 8 shows the structure of the plasma discharge inside and outside of the DBD microplasma channels at different operating pressures. The figure shows luminosity images of the plasma reactor, looking directly into the channels from above (see figure 4). The pink glow is due to the plasma, the bluish glow around it is due to the dielectric (note that the holes in the electrodes have twice the diameter of the channels). Figure 9 shows short-exposure ICCD images of single pulses in one channel under similar conditions.

At atmospheric pressure, figure 8(a), the discharge is concentrated as an annulus on the inner wall of the channel and no discharge is visible in the center of the holes. This is related to the short mean free path of the electrons due to the inelastic collisions with the neutral atoms, which narrows the discharge region. Outside the holes, a micro-filamentary discharge from the hole to the cathode ring can be observed (the white reddish lines traversing the bluish regime). The single-pulse ICCD image in figure 9(a) shows similar behavior.

At a pressure of 100 mbar, figure 8(b), the mean free path of the electrons increases, resulting in a wider and more uniform annular ring inside the holes. Also, note the absence of micro-filamentary discharges in the short-exposure image (figure 9(b)). Meanwhile, the glow discharge extends beyond the confinement of the holes, giving rise to the faint pink glow over the whole field of view. By decreasing the pressure further to 50 mbar, figures 8(c) and 9(c), the plasma discharge is capable of reaching the center and uniformly fills the whole channel.
3.3. Temperature measurements

Gas temperature is a very important parameter in plasma science for two reasons. First, the gas temperature has a direct effect on reactive species generation \[ \text{(25)} \]. Second, there are many applications that need control of thermal behavior, such as wound treatment, chemical decomposition, combustion, etc. Nanosecond plasma discharges typically are of non-equilibrium nature resulting in different kinetic energy distributions for the different species (i.e. electrons, ions, neutrals). Apart from that, the different degrees of freedom (electronic, translation, vibration and rotation) are also not necessarily in equilibrium \[ \text{(26, 27)} \]. In the following sections, we will focus on the rotational and vibrational temperature measurement of the plasma discharge in the DBD microplasma reactor, as well as the average gas temperature.

3.3.1. Rotational temperature

The rotational temperature of the plasma discharge in stagnant air has been obtained by fitting the experimentally observed spectra of the (0–2) band structure of the second positive system (SPS) of \( \text{N}_2 \) (\( C^3\Pi_u \rightarrow B^3\Sigma_g^+ \)) with the Specair simulation tool \[ \text{(28)} \] convoluted with the measured instrumental slit function. The spectra are integrated over 20 ns following the start of the voltage pulse, using a spectrograph slit width of 20 \( \mu \text{m} \). The
3.3.2. Vibrational temperature

The measurements are carried out for a pulse repetition frequency of 3 kHz using the grating of 3600 grooves/mm. The signal has been accumulated over 300,000 pulses to increase the signal-to-noise ratio. The signal was collected from an area covering four complete holes at the center of the reactor disc. The measurements are carried out for a pulse repetition frequency of 3 kHz using the grating of 3600 grooves/mm.

As shown in figure 10, when comparing the rotational line structure with a simulated Boltzmann rotational distribution, a good fit was obtained at both atmospheric pressure and reduced pressure. This implies that the rotational distribution has thermalized (via heavy species collisions or electron collisions) in a short time (few nanoseconds). The rotational distribution of the excited states can be considered representative of the ground state rotational temperature, which is usually close to the gas temperature [29]. The rotational temperatures obtained from the Specair fitting procedure were 550 K ± 30 K and 470 K ± 30 K for pressures of $p = 1$ bar and $p = 50$ mbar, respectively. This reduction in the rotational temperature with pressure can be explained by the reduction in the consumed power, as shown in figure 7(b).

3.3.3. Average gas temperature

To measure the average gas temperature downstream of the reactor, we have used a thermocouple type K (model Testo 0613-1912) with a 5 mm probe diameter and reaction time of 35 seconds to damp the noise. The probe junction is located 5 mm downstream from the central hole of the reactor. Figure 14 shows the air temperature differences were observed between the incident pulse at $t = 10$ ns and the reflected pulse at 50 ns, as can be seen in figure 13. These spectra have been recorded in stagnant air at atmospheric pressure using the 1800 grooves/mm grating centered at 355 nm to cover the (0–1) and (1–2) bands of the SPS of N₂. About 30 percent reduction in emission intensity has been observed between the incident and reflected pulses.
temperature as a function of pulse repetition frequency for an air flow rate of 3.4 l min$^{-1}$ at atmospheric pressure.

As shown in the figure, there is a linear relationship between the pulse repetition frequency and average gas temperature, with a slope of 6.5 K kHz$^{-1}$. From this relationship, we can extract an average value of the thermal power generated by the plasma discharge,

$$Q_{\text{plasma}} = m \cdot c_p \cdot \Delta T,$$

where $m$ is the mass flow rate, $c_p$ is the specific heat of air at constant pressure and $\Delta T$ is the difference between inflow and outflow temperatures. This gives an average value of 0.45 W for 5 kV pulses at a repetition frequency of 1 kHz, or in other words, the energy spent on gas heating is 0.45 mJ/pulse for all channels combined.

This value is 85% of the pulse energy calculated from the voltage and current measurements at atmospheric pressure in section 3.1. The rest of the consumed power is spent on other degrees of freedom such as dissociation of the molecular gas components (such as the dissociation of the molecular oxygen in the quenching reaction of N$_2$(B) [30]), heating of the dielectric material and conduction to the metal body. In the case of reduced pressure, it is expected that the fraction of the power, which is dissipated into gas heating will be lower, and more power will go to the chemical processes due to the higher electron energy and hence ionization efficiency.

3.4. Effective reduced electric field strength ($E/N_{\text{eff}}$)

Determination of the actual reduced electric field ($E/N$) by direct methods is relatively complicated for pulsed micro-discharges because both the electric field ($E$) and the concentration of neutral particles ($N$) vary strongly in time and space. However, Paris et al [31] introduced an indirect
method by using the intensity ratio of nitrogen spectral bands. For this, the (0-0) transition band of the first negative system of N2^+(FNS; B^0Σ_g^+ → X^2Σ_g^+) at 391.4 nm and the SPS (2–5) band at 393.3 nm are used. The following relation between the intensity ratio \( R_{391/394} \) and the reduced electric field has been proposed for low-temperature plasmas in air at a pressure range of 0.3–100 kPa with a maximum deviation of 12%:

\[
R_{391/394} = 46 \exp \left[ -89 \left( \frac{E}{N} \right)^{0.5} \right].
\]

The distribution of the N2^+(B – X) (0–0) and N2 (C – B) (2–5) intensities over the channel width are presented in figure 15, at 300 and 50 mbar, for both the incident and reflected pulses at 5 kV and 3 kHz pulse repetition rate. The spectra of the N2^+(B – X) (0–0) and N2 (C – B) (2–5) bands are accumulated for 10 ns after the onset of each pulse and integrated over 300,000 pulses using a 1800 grooves/mm grating.

Three distinct regimes can be recognized from these figures when going from the center of a channel outwards: (1) inside the channel (A-B), (2) between the channel and the grounded electrode edge (B-C), and (3) the grounded electrode (C-D). At 300 mbar, and during the incident pulse, the minimum intensity of the nitrogen FNS is located at the center of the channel (point A). Going outwards, two peaks can be distinguished. The first peak is located inside the channel close to the wall (point B). This is caused by the integration over the line of sight of the ionization wave, which is traveling from the high-voltage electrode towards the channel exit. This peak is expanding by reducing the pressure until it reaches the center at 50 mbar when the central minimum disappears.

The second peak is located at the ground electrode edge (point C), which is most likely produced by a discharge, which is formed due to the field enhancement at the ground electrode edge. The intensities of the two peaks have almost the same strength, even at different operating pressures. In contrast, during the reflected pulse, only one peak of N2^+(B – X) (0–0) is formed, the one at the channel walls, while the one at the grounded edge is absent. This could be explained by the following three effects. First, the high electron density left over from the incident pulse around the grounded electrode covers the sharp edge, which consequently reduces the local field enhancement as well as the energy and number of produced electrons. Second, it could be related to the drift direction of the electrons. Third, the low voltage of the reflected pulse compared to the incident pulse may prevent the N2^+(B – X) (0–0) excitation.

The vibrationally excited nitrogen state N2 (C – B) (2–5) appears mostly near the wall of the channel during both the incident and reflected pulses, although with about half of the intensity during the reflected pulse. Furthermore, it seems like the discharge at the grounded electrode, near point (C), consumes most of the energy on the ionization and less on vibrational excitation. This can explain the reduced amount of N2 (C – B) (2–5) compared to N2^+(B – X) (0–0) at this location.

For pressures between one atmosphere and 100 mbar, the discharges have almost identical structures and only differ in the relative intensities between the excited states. However, at 50 mbar, the structure changes dramatically, as shown in the bottom graphs in figure 15. The major difference in the structure is that there is only one peak in the center of the channel for both the SPS of N2 and FNS of N2^+. This is due to the large main free path at 50 mbar, as discussed earlier in
section 3.2. It is also clear that the FNS of $N_2^+$ is about one order of magnitude higher than the SPS of $N_2$, which indicates a high reduced electric field strength.

Figure 16 shows the effective reduced electric field strength $(E/N)_{\text{eff}}$, which is determined by the intensity ratio method, presented in equation (3), for the incident and reflected pulses. A spatial averaging has been performed for the $N_2^+ (B \rightarrow X) (0 \rightarrow 0)$ and $N_2 (C \rightarrow B) (2 \rightarrow 5)$ over (A-B) and (B-D) positions for one channel. The signals are accumulated for 10 ns and integrated over the line of sight for 300,000 pulses.

The general trend, for all lines in the figure, is that the effective reduced electric field strength increases gradually with decreasing operating pressure. This trend is expected, as the neutral number density of the gas decreases with pressure, while the applied voltage (and therefore the path-integrated electric field) remains constant. The figure shows a value of $550 \pm 100$ Td and $340 \pm 50$ Td for $(E/N)_{\text{eff}}$ for the incident pulse inside the channel (A-B) and outside the channel (B-D), respectively. By reducing the pressure to 100 mbar, $(E/N)_{\text{eff}}$ increases gradually to $4600 \pm 700$ Td and $1900 \pm 300$ Td. Also, at this pressure range, $(E/N)_{\text{eff}}$ for the reflected pulse inside the channel is slightly higher than for the incident pulse. This can be explained by the substantial increase in $(E/N)_{\text{eff}}$ of the incident pulse outside the channel. By reducing the pressure down to 50 mbar, $(E/N)_{\text{eff}}$ inside the channel increases to about $10000 \pm 1500$ Td for the incident pulse with a relatively lower value outside the channel.

The residence time of a volume of gas passing through the discharge region in a channel (length of 0.36 mm for single-layer operation) ranges from 50–250 $\mu$s for air velocities between 7–1.4 m s$^{-1}$. Obviously, the minimum residence time is much longer than the characteristic time scale of the plasma discharge, which is on a tens of nanosecond time scale. However, these residence times are slightly shorter than the inter-pulse delay at a pulse repetition frequency of 3 kHz. This can explain the reduction in emission intensity at higher flow velocities, as shown in figure 17. The figure shows a roughly 25% reduction in emission intensity when increasing the flow from 0 to 1.4 m s$^{-1}$ and a smaller, more gradual, reduction when increasing the flow further up to 7 m s$^{-1}$. In stagnant air, all high-voltage pulses ionize the same volume of gas in the discharge region. This increases the ionization level before the new breakdown occurs, which leads to higher emission intensity [32, 33], while at higher velocities most of the gap is filled with fresh gas and less pre-ionization will exist.

3.6. Reactor lifetime assessment

One of the serious problems of microplasma devices is their often short lifetime due to surface and internal damage. This is due to the frequent loss of plasma confinement due to vaporization and ablation, which cause significant surface erosion and structural failure [34, 35]. Reactor lifetime is very sensitive to many parameters, e.g. geometry, material of the dielectric and power source.

Figure 18 shows the dependence of the reactor lifetime on the pulse repetition frequency at atmospheric pressure and 50 mbar. At a frequency of 2 kHz and atmospheric pressure, the reactor lifetime was about 24 min in continuous operation mode. While at 50 mbar, it was about 100 min. This shows an enhancement in the reactor lifetime of about four times by lowering the operating pressure by a factor of 20. This enhancement increases to a factor of about ten at a pulse repetition rate of 10 kHz, even though the lifetime itself decreases in both cases.
Two factors may contribute to the observed enhancement. First, at low pressure, the ion impact frequency is low, which slows down the dielectric wear-down process. Second, at low pressure the gas temperature is lower, as was shown in section 3.3.1. Both reasons make the reactor more prone to failure at atmospheric pressure due to thermal stress on the internal structure of the dielectric material. Other factors that may affect the lifetime of the reactor and need more investigation are voltage amplitude and duration, geometry, reactor material and operating gas.

4. Summary and conclusion

In this work, a new DBD microplasma flow reactor driven by nanosecond high-voltage pulses has been presented. With this design we were able to sustain a non-equilibrium plasma discharge in a pattern of 400 μm diameter channels at pressures up to 1 bar. The DBD microplasma flow reactor is a promising device for a wide range of applications that require an efficient interaction between a non-thermal plasma discharge and a gas flow. Some of these applications are pollutant control, CO₂ to CO conversion, plasma-assisted combustion and plasma medicine.

Time-resolved electrical and optical measurements have been conducted to characterize the main features of the plasma discharge in the DBD microplasma reactor. Pulse energies per channel of about 1.46 and 1.3 μJ have been calculated from the current and voltage time evolution at atmospheric pressure and 50 mbar, respectively. The discharge at low pressure is characterized by high vibrational temperatures (roughly 4000 K) and high effective reduced electric field strengths (10000 Td) compared to atmospheric pressure (3460 K and 550 Td), which indicates a higher electron energy at lower pressure. Based on luminosity images, all gas flowing through the reactor can be assumed to interact with the plasma at 50 mbar. At higher pressures, the emission concentrates at the channel walls and not all gas is in direct contact with the plasma. The distribution of the plasma-generated radicals, however, is unknown. The luminous part of the plasma is strongly non-thermal, with vibrational temperatures of about 4000 K, but the gas temperature remains only a few degrees above ambient.

In addition, we have noticed that the discharge emission intensity decreases slightly with increasing air flow velocity through the channels at atmospheric pressure due to the lower pre-ionization level at higher velocities. Finally, a reactor lifetime study showed around 100 min lifetime at low pressure. Nevertheless, more effort is needed to increase the lifetime of the reactor in order to transfer it to real applications.

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