A Novel Pulsed Corona Discharge Reactor Based on Surface Streamers for NO Conversion from N2-O2 Mixture Gases

M. A. Malik  
*Old Dominion University*

Karl H. Schoenbach  
*Old Dominion University*

Follow this and additional works at: [http://digitalcommons.odu.edu/bioelectrics_pubs](http://digitalcommons.odu.edu/bioelectrics_pubs)

Part of the [Biomedical Engineering and Bioengineering Commons](http://digitalcommons.odu.edu/bioelectrics_pubs) and the [Environmental Sciences Commons](http://digitalcommons.odu.edu/bioelectrics_pubs)

Repository Citation  
[http://digitalcommons.odu.edu/bioelectrics_pubs/36](http://digitalcommons.odu.edu/bioelectrics_pubs/36)

Original Publication Citation  

This Article is brought to you for free and open access by the Frank Reidy Research Center for Bioelectrics at ODU Digital Commons. It has been accepted for inclusion in Bioelectrics Publications by an authorized administrator of ODU Digital Commons. For more information, please contact digitalcommons@odu.edu.
A Novel Pulsed Corona Discharge Reactor Based on Surface Streamers for NO Conversion from N₂-O₂ Mixture Gases

M. A. Malik and K. H. Schoenbach
Frank Reidy Research Center for Bioelectrics, Old Dominion University, USA

Abstract—A novel pulsed corona discharge reactor is described which utilizes surface-plasma along insulating surfaces. The electrodes are comprised of a stainless steel wire anode of 150 µm diameter stretched along the surface of a glass sheet and two parallel aluminum strips as cathodes. An eight-stage Marx bank, which provides 60 ns, 40–45 kV monopolar pulses, was used to produce the surface streamers in nitrogen-oxygen mixtures at atmospheric pressure. With increasing oxygen content, the energy efficiency for NO₂ and O₃ synthesis was found to increase. The energy efficiency is almost the same for the surface-plasma and volume-plasma. However, the surface-plasma was found to be significantly more energy efficient than the volume-plasma for conversion of dilute NO in a feed gas containing 0–15% oxygen and with the balance being nitrogen. It is explained on the basis of surface-mediated reactions, the electric wind effect, and the diffusivity of the plasma which covers a larger fraction of the volume of the discharge gap as compared to volume-plasma. The surface-plasma reactor will be used to explore the treatment of NOₓ and hydrocarbons in diesel engine exhaust.

Keywords—NOₓ conversion, ozone synthesis, nitrogen fixation, non-equilibrium plasma, surface flashover

I. INTRODUCTION

Streamers along insulating surfaces have been studied mainly for the purpose of determining the dielectric strength of insulators in high voltage systems [1, 2]. Modeling results show that the surface streamers consist of a positive streamer head followed by quasi-neutral plasma in the channel behind the head and surrounded by a layer of positive charges [3]. This model explains the measured positive residual charge on the surface. Experimental results also indicate that the insulating surface plays a major role in the development of the surface-plasma. For example, photo-electron emission from the surface is believed to increase the ionization rate, which explains the faster speeds of surface streamers as compared to streamers in the gas alone [1, 2].

Interest in the application of surface-plasmas for environmental pollution control is growing because of their better energy efficiency as compared to the corresponding volume-plasmas [4–7]. For example, conversion of NO [4], destruction of toxic VOCs [5] and synthesis of O₃ [6] from air have been compared in surface-plasma and volume-plasma in a dielectric barrier discharge reactor all showing better energy efficiency with surface-plasma as compared to volume-plasma. A significant improvement in energy efficiency for destruction of toxic VOCs in a surface-plasma as compared to the volume-plasma in a pulsed corona discharge reactor has been reported [7]. Conversion of NO from N₂ + O₂ mixtures in the surface-plasma and the corresponding volume-plasma in a pulsed corona discharge reactor is being reported in this manuscript.

II. EXPERIMENTAL

The experimental setup employed in this study is shown in Fig. 1. The pulsed power system comprises a DC high voltage power supply and an 8-stage Marx bank. The total capacitance (C) of the Marx bank after switching is 0.29 nF, and the inductance (L) of the circuit is 1.53 µH. In order to obtain monopolar pulses, the resistive load (R) must exceed 2 (L/C)⁽¹/₂⁾. The shortest possible monopolar pulse duration for an LCR circuit is obtained when R = 2 (L/C)⁽¹/₂⁾ (critically damped pulse). For the Marx circuit, the resistance has been calculated as R = 146 Ω. This is a value that is small compared to the resistance of the streamer discharge. Therefore, placing two resistors of 75 Ω each in parallel to the discharge guarantees that the voltage pulse applied to the discharge is always the same, even when the discharge resistance fluctuates.

Voltage and current waveforms were recorded on an oscilloscope (Tektronix TDS 3052). The voltage probe was a Tektronix P6015A with a bandwidth of 75 MHz. Two 75 Ω resistors in series, both parallel to the load, were utilized as a 1:1 voltage divider. The current probe was a Pearson Electronics Current Monitor with a sensitivity of 0.1 V/A, and bandwidth of 20 MHz (Model 110A). The temporal resolution of the voltage and current traces is dependent on the bandwidth of the lowest bandwidth element in the diagnostic circuit, which in our case was the Pearson current probe. For consistency we have chosen a bandwidth of 20 MHz, by using a 20 MHz filter in the oscilloscope.

The electrical power was calculated as the product (VI) of the measured pulse voltage (V) and current (I). The energy per pulse (Eₚ) is the time integral (∫VI dt) of the power for the duration of the pulse. For each experiment, Eₚ was calculated from an average of 64 pulses and pulse frequency (f) from an average of 40 experiment.
Fig. 1. Experimental setup: DC high voltage power supply connected to an 8-stage Marx bank and reactor. 1 = 100 kΩ resistors, 2 = 2.3 nF capacitors, 3 = spark gaps, 4 = 75 Ω resistors, 5 = reactor, 6 = high voltage probe, 7 = current probe, 8 = oscilloscope, 9 = NOx analyzer, 10 = gas chromatograph, 11 = gas bottles, 12 = needle valves, and 13 = flow meters.

Fig. 2. Surface-plasma reactor: a) is a partially expanded view, and b) is view of the assembled reactor, and c) shows a time integrated image of the surface-plasma in nitrogen generated by a single voltage pulse. The components are: 1 and 3 are top and bottom glass sheets; 4 is the stainless steel wire anode of 150 µm diameter; 2 is a spacer comprising Plexiglas or Teflon end fittings and cathodes (number 5) of aluminum forming two sides of the spacer, and 6 are gas inlet/outlet. The whole assembly is enclosed in two Plexiglas sheets held together by nuts and bolts and sealed by silicon sealant.

Fig. 3. Volume-plasma reactor: a) is a sketch of the reactor where the components are: 1 is a Plexiglas cylinder of 4.5 cm ID, 5.1 cm OD, and 15 cm length; 2 are Plexiglas end-fittings; 3 is a cylindrical cathode of 4.5 cm OD, 10 cm length, made of stainless steel mesh; 4 is a wire anode made of stainless steel wire of 150 µm diameter and stretched along the axis of the cylinder; and 5 are gas inlet/outlet; b) shows a time integrated image of volume-plasma generated in nitrogen by a single voltage pulse.

Gases, i.e., N₂, O₂ and NO were supplied from high pressure gas bottles. Flow rates of the gases were controlled by needle valves and monitored with ball-float flow meters. A mixture of N₂ and O₂ with or without NO was passed through the reactor at a rate of 1 liter per minute (L/min) at a temperature of 25°C and one atmosphere of pressure in all experiments. The initial concentration of NO was 300 ppm in surface-plasma and plasma reactor in a wire-to-cylinder arrangement of electrodes and a time-integrated image of the volume-plasma.

Fig. 2 shows the schematics of the surface-plasma reactor and a time-integrated image of the surface-plasma. The reactor comprises a wire to two parallel-plate electrodes stretched on the surface of a dielectric sheet and enclosed by another dielectric sheet with a spacer in between. The electrode lengths, inter-electrode gaps and thickness of the spacer that defines the volume of the discharge gap are listed in the results and discussion section. Fig. 3 shows the schematics of the volume-plasma reactor in a wire-to-cylinder arrangement of electrodes and a time-integrated image of the volume-plasma.

The pulse frequency (f) was ~10 Hz in all experiments.
700 ppm in volume-plasma in all experiments, except where mentioned otherwise. The concentrations of O$_2$, NO, and NO$_2$ were monitored by a NO$_x$ analyzer (ENERAC Model 500, equipped with O$_2$, NO and NO$_2$ sensors). Time-integrated images of single plasma discharges were recorded using a 5 mega-pixel camera.

### III. RESULTS AND DISCUSSION

Fig. 4 illustrates two geometries of electrodes and dielectric surfaces usually employed for the study of surface-plasmas [2]. The geometry illustrated in Fig. 4a is similar to the geometry utilized in dielectric barrier discharges and the one illustrated in Fig. 4b is similar to that used for corona discharges, such as pulsed corona discharges. The latter is employed in the present study.

Two plasma reactors, i.e., a surface-plasma reactor (Fig. 2) and a volume-plasma reactor (Fig. 3) are employed in this study. The placement of electrodes on the dielectric surface ensured maximized surface-plasma formation in the surface-plasma reactor. In the case of the volume-plasma reactor, the cylinder electrode was kept 2.5 cm away from the end-fittings, which minimized the probability for the generation of surface-plasma due to the increased distance between the electrodes along the end-fitting surfaces. Therefore the reactor shown in Fig. 3 is optimized for volume-plasma.

Comparison of time integrated images of the plasmas show that the surface-plasma channels shown in Fig. 2c are thicker, more diffuse, and cover a larger fraction of the discharge gap compared with the volume-plasma channels shown in Fig. 3b, under the same experimental conditions.

The dielectric surface is likely to be charged positively [3] by the ions in the streamer. The resulting positive space charge causes a repulsive force on the positive surface charges surrounding the streamer. Since the expansion of the streamer towards the opposite dielectric surface is restricted to its positive surface space charge, the repulsive forces may lead to an expansion perpendicular to the streamer axis in the space between dielectric sheets rather than a shift away from the dielectric surface, and consequently, to a more diffuse streamer. This qualitative explanation would need to be confirmed by modeling.

Typical voltage and current waveforms observed in surface-plasma reactor are illustrated in Fig. 5. In the range of our experiments, changes in the ratio of oxygen and nitrogen concentrations did not seem to influence the pulse shapes significantly. The peak voltage was 40-45 kV, pulse rise time was ~15 ns, and pulse width at half maximum was ~60 ns.

The current reached a maximum value of ~16 A at ~25 ns and at a voltage of ~35 kV, still in the rising part of the voltage pulse. Visual observation and images of the discharge confirmed bridging of the electrode gap by
surface streamers. We believe that a streamer-to-glow transition takes place after the streamers bridge the gap between the electrodes. The measured resistance at the peak current is \( \sim 2 \, \Omega \), far above values for arcs. However, the point of transition from streamer to glow discharge could not be determined from the temporal development of voltage and current waveforms.

The electrical energy is mainly used for electron heating rather than gas heating in the plasma generated by high voltage pulses of sharp rise time and short duration [8]. High-energy electrons generated in the plasma collide with ambient gas molecules and dissociate them into free radicals, like O and N [9]. The radicals then react with each other and with ambient gas molecules and produce new compounds like NO, NO\(_2\), O\(_3\), etc., by reactions such as the following:

\[
\begin{align*}
    O_2 + e^* & \rightarrow O + O + e \quad (1) \\
    N_2 + e^* & \rightarrow N + N + e \quad (2) \\
    N_2^* + O_2 & \rightarrow O + O + N_2 \quad (3) \\
    O + O_3 + M & \rightarrow O_3 + M \quad (4) \\
    \text{where } M \text{ is } O_2 \text{ or } N_2 \\
    O_2^* + O_3 & \rightarrow 2O_2 + O \quad (5) \\
    N_2^* + O + N & \rightarrow NO + N_2 \quad (6) \\
    N^* + O + N_2 & \rightarrow NO + N_2 \quad (7) \\
    N^* + NO & \rightarrow N_2 + O \quad (8) \\
    NO + NO + O_2 & \rightarrow 2NO_2 \quad (9) \\
    NO + O_3 & \rightarrow NO_2 + O_2 \quad (10) \\
    NO + O + M & \rightarrow NO_2 + M \quad (11)
\end{align*}
\]

The concentration of NO\(_2\) in the treated gas indicates the amount of nitrogen- as well as oxygen-based active species produced in the plasma. NO\(_2\) was measured in parts per million (ppm), volume-to-volume, in exhaust gas at 25°C and one atmospheric pressure after three minutes of plasma operation at 10 Hz. The energy cost (\( \epsilon \)) in units of electron-volts per molecule (eV/molecule) was calculated using the formula: \( \epsilon = 250 \frac{E_g/(F\Delta NO_2)}{F} \), where \( F \) is the flow rate in liters per second (L/s) and \( \Delta NO_2 \) is the change in concentration of NO\(_2\) in ppm [10]. The energy yield (\( Y \)) in units of gram per kilowatt-hours (g/kWh) is calculated by using the formula: \( Y = 37.3 \frac{M}{\epsilon} \), where \( M \) is molecular weight of the compound. It should be mentioned here that “energy efficiency” in this manuscript refers to energy yield (\( Y \)) and it is reciprocal of energy cost (\( \epsilon \)).

Fig. 6 shows the energy cost for NO\(_2\) production as a function of oxygen concentration in ambient gas in the volume-plasma reactor and two surface-plasma reactors: one having spacer thickness 1.4 cm, and the other 0.2 cm. The energy cost gradually decreased from \( \sim 30 \) eV/NO\(_2\) molecule (48 g/kWh) at 3% oxygen to \( \sim 15 \) eV/NO\(_2\) molecule (96 g/kWh) in air (20.9% oxygen) in all three reactors, in agreement with earlier studies [11]. The energy efficiency in volume-plasma and the surface-plasma reactors measured in this study is almost the same. There was no NO detected in the exhaust gas. Ozone formed in the plasma reactor converts NO to NO\(_2\) by reaction No. 10 [12], which explains the absence of NO in the treated gas. Ozone reacts preferentially with NO, but in the absence of NO it can oxidize NO\(_2\) to higher oxidation states, such as NO\(_3\) and N\(_2\)O\(_5\) [12]. The oxides of nitrogen other than NO and NO\(_2\) were not analyzed in this study.

In the present study, excess NO was mixed with the plasma exhaust and the reduction in the amount of NO was monitored. Energy cost for ozone in the plasma exhaust was calculated by assuming one molecule of ozone oxidized one molecule of NO to NO\(_2\). Fig. 7 shows that the energy cost for ozone production gradually decreased with an increase in oxygen content from \( \sim 45 \) eV/molecule (\( \sim 34 \) g/kWh) at 3% oxygen to \( \sim 13 \) eV/molecule (\( \sim 116 \) g/kWh) at 20.9% oxygen. Otherwise, the energy cost for ozone production was similar in the three reactors tested in this study.
Fig. 7. Energy cost ($\varepsilon$) for O$_3$ production versus oxygen concentration in the feed gas under three different conditions:

-○-- volume-plasma reactor, ----×---- surface plasma with spacer thickness 1.4 cm, and ─∆─ surface plasma with spacer thickness 0.2 cm (the electrode lengths were 30 cm and the gaps between anode and cathode were 4.5 cm in both the surface plasma reactors).

Fig. 8. Energy cost ($\varepsilon$) for NO conversion versus oxygen concentration in feed gas under three different conditions:

-○-- volume-plasma reactor, ----×---- surface plasma with spacer thickness 1.4 cm, and ─∆─ surface plasma with spacer thickness 0.2 cm (the electrode lengths were 30 cm and the gaps between anode and cathode were 4.5 cm in both the surface plasma reactors).

A major fraction of NO$_x$ in flue gas is NO. The NO is relatively difficult to remove from the flue gas compared to higher oxidation states of nitrogen, such as NO$_2$. The NO$_2$ is easier to remove, e.g., by scrubbing and chemical treatment [13]. This technique may be successful for NO$_x$ treatment from stationary sources, such as coal or gas fired power plants. However, it is less likely to be applied in the case of mobile sources, such as diesel engine exhausts in vehicles, due to space limitations. Direct treatment by plasmas followed by hydrocarbon selective catalytic reduction (H-SCR) is being developed for treating NO$_x$ from the diesel engine exhausts [14, 15]. In the direct treatment, the NO is oxidized to NO$_2$ and, at the same time, any unburned hydrocarbons in the exhaust are activated by partial oxidation.

A synthetic mixture of NO + N$_2$ + O$_2$ was fed to the reactors for the direct plasma treatment. Fig. 8 shows the conversion of NO fed to the reactors. The production of NO$_2$ is not shown because when a synthetic mixture of NO and NO$_2$ (balance N$_2$) was analyzed, the accuracy of NO estimation was good (2% of the value), but accuracy of NO$_2$ was not. Further, NO$_2$ concentration exceeded the analyzable limits of 500 ppm in most of the experiments.

The energy cost for NO conversion was found to increase with a decrease in oxygen content from 20.9% to 3% and showed a sudden decrease in the case of pure nitrogen (0% oxygen) in the ambient gas in all the
reactors. These trends are in agreement with earlier studies [16].

The NO is removed either by reduction by N (reaction No. 8) or by oxidation by O or O₃ (reaction No. 10 & 11). The reduction by N is counterbalanced by NO formation (reaction No. 7). So, there is a net oxidation reaction responsible for NO removal [9]. The role of the oxidation reaction is further supported by the fact that the bond dissociation energy of O₂ (5.1 eV) is smaller than that of N₂ (9.8 eV). With a decrease in oxygen content in the ambient gas, oxidizing species like O and O₃ decrease. This explains the increase in energy cost for NO removal. In the extreme case of pure nitrogen, NO is removed by reduction with N, while the NO formation by N* becomes limited due to the absence of oxygen. It explains the low energy cost for NO removal in pure nitrogen as compared with the case of 3% oxygen mixed with nitrogen.

Fig. 8 shows that the surface-plasma is more energy efficient (lower energy cost) compared to the volume-plasma when oxygen in the ambient gas is less than that in air. Diesel exhaust contains oxygen in the range of 3-15%. The presence of unburned hydrocarbons in the case of real diesel engine exhaust will further increase the NO conversion significantly [9, 17]. These facts clearly indicate the advantage of using surface-plasma for treating diesel exhaust.

The energy cost for nitrogen-based and oxygen-based reactive species are expected to be almost equal for surface-plasma and volume-plasma, since the energy cost for their end products, i.e., NO₂ and O₃ are almost equal. So, the production of free radicals is less likely the factor responsible for the difference in the energy cost for NO conversions shown in Fig. 8. Nitrogen and oxygen molecules, being abundant in the treated gas, are available in sufficient concentration inside the plasma. So the reactive species do not need to enter the gas surrounding the plasma zone for production of NO₂ and ozone. NO, on the other hand, is in very low concentration and the reactive species formed in the plasma need to extend into the ambient gas. The diffusible nature of the plasma covering a larger fraction of the reaction zone in the case of surface streamers most probably extends the reach of the reactive species as compared with volume streamers.

Electric wind generated by the discharge is another factor considered responsible for efficient mixing and, consequently, energy efficient chemical conversion in surface dielectric barrier discharges [5]. Adsorption and stabilization of reactive species, such as, O [18] and N [19] from plasma onto glass and their availability for further oxidation reactions, such as oxidation of NO to NO₂ by reacting with adsorbed O [18] has been shown in recent literature. The surface-mediated reactions can also explain better energy efficiency in surface-plasmas compared to the volume-plasmas. The dielectric surfaces in the case of surface-plasma have a higher probability of adsorbing and stabilizing the reactive species due to their close proximity with the plasma compared to the case of volume-plasma. These three factors, i.e., diffused plasma, electric wind, and surface mediated reactions, explain the lower energy cost for NO conversion in the surface-plasma than in the volume-plasma.

One exception to the general trend mentioned above is that the volume-plasma was more energy efficient for NO conversion from air (20.9% oxygen) than the surface-plasma. The reason for this reversal of the trend in air is not understood at this stage.

The energy per pulse was ~100 ±10 mJ in the case of volume-plasma and it was ~35 ±8 mJ in the case of surface-plasma employed for data shown in Figs. 6, 7 and 8. The energy cost for NO₂ and O₃ synthesis (Figs. 6 and 7) was almost the same, while the energy cost for NO conversion (Fig. 8) is slightly lower in surface-plasma in which the spacer thickness was 0.2 cm compared to the surface-plasma having space thickness of 1.4 cm. It means that the volume of the discharge gap of the surface-plasma reactor can be reduced from ~400 ml to ~50 ml by reducing the thickness of the spacer from 1.4 cm to 0.2 cm without reducing the energy efficiency.

Fig. 9 shows the dependency of energy cost on the degree of NO conversion in the volume-plasma and surface-plasma formed on Teflon or Plexiglas or glass dielectric. Energy per pulse was ~100 ±10 mJ in volume-plasma and ~10 ±2 mJ in surface-plasma employed for NO conversion shown in Fig. 9. In these experiments, the inlet concentration of NO was varied, which resulted in different degrees of NO conversions, keeping all other parameters the same. The energy cost increases with the degree of NO conversion. This is a general trend in any plasma treatment. The energy cost for surface plasma on glass, poly(methyl methacrylate) and teflon as dielectric layer is significantly lower compared to the volume plasma. For example, the energy cost at 50% NO conversion is 175 eV/molecule (6 g/kWh) in the case of volume plasma and is reduced in surface plasma to 97 eV/molecule (12 g/kWh) on Teflon, 74 eV/molecule (15 g/kWh) on poly(methyl methacrylate), and 51 eV/molecule (22 g/kWh) on glass.

Fig. 9 shows the order of energy cost with respect to the dielectric surface: Teflon > poly(methyl methacrylate) > glass. This trend indicates that the surface plasma strongly interacts with the dielectric surface. This is supported by the fact that the physical characteristics of the plasma are also dependent on the dielectric [1, 2]. Adsorption and stabilization of plasma-produced reactive species on quartz surfaces and their utilization in surface mediated chemical reactions has been reported [18, 19]. The glass surface employed in this study is closer to that of quartz and, consequently, more efficient for the surface mediated reactions, as compared to organic polymers like Teflon or poly(methyl methacrylate). The role of other factors, such as inter-electrode gap, effective length of the electrodes, applied voltage, pulse frequency and other dielectrics, especially porous ceramics having larger surface area, needs to be evaluated.

The energy efficiency for 50% NO conversion from nitrogen by volume-plasma reactor in this study (~6
g/kWh) is the same as calculated from the data shown in earlier studies on pulsed corona discharges and dielectric barrier discharges [20]. It is comparable with ~5 g/kWh to that of semi-wet type pulsed corona discharges [21]. The energy efficiency was increased to about 22 g/kWh by employing surface-plasma in the present study. It should be mentioned that the pulsed corona discharge reactor employed in this study is not the optimum system for this application. Its energy efficiency can be improved further by additives and by varying electrical parameters. For example, additives, such as hydrocarbons [17] or ammonia [22] can significantly improve the energy efficiency for NO conversion in plasma reactors. Energy efficiency value of 125 g/kWh has been reported in a plasma reactor with ammonia addition [22]. A good example of energy efficiency improvement by varying electrical parameters is ~18 g/kWh energy efficiency for 50% NO conversion in a pulsed corona discharge reactor biased by very short 5 ns duration pulses [23]. The best energy efficiency in the present study was ~22 g/kWh achieved by employing the surface-plasma reactor.

5. The energy cost for the plasma chemical reactions, such as NO conversion, is dependent on the dielectric surfaces in the case of surface plasma.

ACKNOWLEDGMENT

This work is supported by the “Frank Reidy Fellowship in Environmental Plasma Research.” The authors thank J. Thomas Camp for providing the Marx Bank and for helpful discussions related to the high voltage system. The authors thank Shu Xiao and Juergen F. Kolb for useful technical discussions. The authors thank Mr. Kevin Sullivan from ENERAC Inc. for re-checking cross-sensitivity of NO and NO₂ sensors when a mixture of NO and NO₂ (balance N₂) is fed to the NOₓ analyzer employed in this study.

REFERENCES


