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
Optimization of Ultraviolet Emission and Chemical Species Generation from a Pulsed Dielectric Barrier Discharge at Atmospheric Pressure

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Optimization of ultraviolet emission and chemical species generation from a pulsed dielectric barrier discharge at atmospheric pressure

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One of the attractive features of nonthermal atmospheric pressure plasmas is the ability to achieve enhanced gas phase chemistry without the need for elevated gas temperatures. This attractive characteristic recently led to their extensive use in applications that require low temperatures, such as material processing and biomedical applications. The agents responsible for the efficient plasma reactivity are the ultraviolet (UV) photons and the chemically reactive species. In this paper, in order to optimize the UV radiation and reactive species generation efficiency, the plasma was generated by a dielectric barrier discharge driven by unipolar submicrosecond square pulses. To keep the discharge diffuse and to maintain low operating temperatures, helium (He) was used as a carrier gas. Mixed with helium, varying amounts of nitrogen (N_2) with the presence of trace amounts of air were used. The gas temperature was determined to be about 350 K at a 1-kHz pulse frequency for all cases and only slightly increased with frequency. The UV emission power density, P_{UV} , reached its highest level when 5% to 10% of N_2 is mixed to a balance of He. A maximum P_{UV} of about 0.8 mW/cm² at 10-kHz pulse frequency for a He(90%)+ N_2 (10%) mixture was measured. This was more than four times higher than that when He or N_2 alone was used. Furthermore, the emission spectra showed that most of the UV was emitted by excited NO radicals, where the oxygen atoms came from residual trace amounts of air. In addition to NO, NO_2 , and excited N_2 , N_2^+ , OH, and He were also present in the plasma. © 2005 American Institute of Physics. [DOI: 10.1063/1.1980530]

I. INTRODUCTION

Atmospheric pressure nonthermal plasmas have recently received increased attention because of several emerging applications such as surface modification of polymers, biological and chemical decontamination of media, aerodynamic drag reduction, and shock-wave mitigation.¹⁻⁶

The main agents that are responsible for the efficient plasma reactivity are ultraviolet (UV) photons and/or the reactive neutral species, and possibly the charged particles.^{7,8} To enhance the ionization and excitation processes, fast-rising voltage pulses (pulse widths from tens to hundreds of nanoseconds range) were suggested by several researchers. Liu and Neiger^{9,10} and Laroussi *et al.*¹¹ applied submicrosecond unipolar pulses with success to enhance energy-transfer efficiency to dielectric barrier discharges (DBD) at low pressure and high pressure, respectively. Liu and Neiger⁹ found that the energy efficiency for ozone synthesis is 30% higher with pulsed DBD than that driven with sine-wave voltages. Mildren and Carman¹² and Carman and Mildren¹³ found that the UV power output from a xenon excimer lamp driven by ~150-ns short pulses had an energy efficiency three times higher than the same lamp excited by sine-wave voltages.

One reason for the better energy transfer to the plasma is that two consecutive discharges occur for each applied voltage pulse. The first discharge occurs at the voltage pulse rising edge, followed by a second discharge at the falling edge of the voltage pulse.⁹⁻¹¹ The second discharge ignites without energy input from the external circuit. The energy

needed is provided by the accumulated surface charges left from the first discharge. These double discharges do not occur under normal low-frequency sine-wave excitation. This process ultimately leads to a much improved power transfer to the plasma.

In this paper, a specially designed pulsed DBD-based discharge was used to study the UV emission and active species generation efficiency with gas mixtures containing different percentages of He and N_2 . The operating pressure was 1 atm, with residual trace amounts of air present in the discharge chamber. First, the gas temperature was measured by comparing the experimental $N_2(C-B)$ 0-0 transition spectrum with simulation results. This was followed by a detailed study of the UV emission power density from the discharge for different percentages of He and N_2 and different pulse frequencies. In Sec. III C, to identify the excited species responsible for the UV emission and other chemically important species, emission spectroscopy was used. Because of the presence of oxygen (from residual traces of air), molecules such as NO and NO_2 were formed. Therefore, a calibrated multigas detector was used to measure the absolute concentrations of NO and NO_2 for different pulse frequencies.

II. EXPERIMENTAL SETUP

Figure 1 is a schematic of the discharge setup. It comprises two parallel electrodes separated by a gap. One of the electrodes is made of a 2 × 2 in² aluminum plate covered by an alumina (Al_2O_3) sheet. The dimensions of the alumina sheet are 75 × 75 × 1 mm³.³ The second electrode is made of a copper disk (diameter of 5.7 cm) with several holes through which the operating gas is injected. The diameter of

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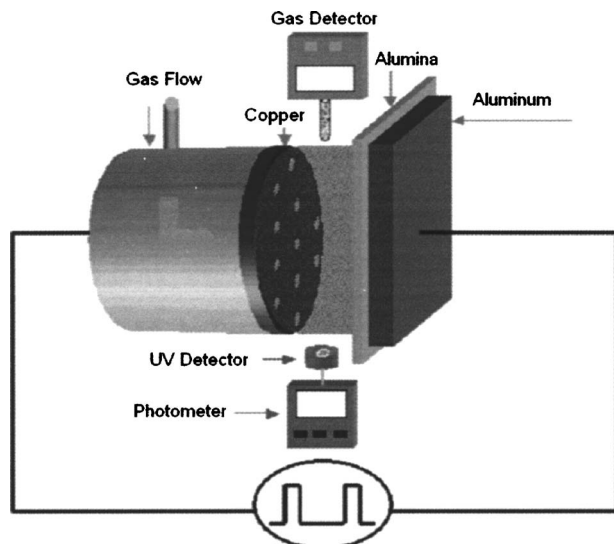


FIG. 1. Experimental setup of the discharge system with diagnostics.

the holes is about 1 mm. The distance between nearby holes is 5 mm. This copper electrode is not covered by a dielectric. The distance of the gap between the electrodes is adjustable from 1 mm to a few centimeters. The gas flows out of the holes and into the discharge gap. A multigas manifold with flow meters was used to adjust the percentage of He and N₂. The plasma was driven by a pulse generator capable of producing narrow high-voltage pulses (pulse width at a few hundred nanoseconds) at adjustable repetition rates. Detailed description of the pulsed power supply can be found in Ref. 11.

For the identification of excited species and the determination of gas temperature, optical emission spectroscopy (OES) was conducted with a half-meter spectrometer (Acton Research SpectraPro 500i). To carry out absolute measurements of the emitted UV power, a radiometer/photometer (International Light, Inc. model: IL1400A) (165–310 nm) was used. The distance from the UV detector to the axis of the copper disk was 6 cm. To identify the excited species responsible for the UV emission and the other chemically important species present in the plasma, emission spectra in the 200–800-nm range were recorded. The absolute concentrations of NO and NO₂ were measured by a multigas detector (RAE System: model PGM-7840). The gas detector was located 1 cm from the plasma surface.

III. EXPERIMENTAL RESULTS AND OBSERVATIONS

For all the experimental results presented in this paper the applied voltage, pulse width, pulse rise time, and the gap distance were $V_a=9$ kV, $t_{PW}=500$ ns, $\tau_r=50$ ns, and $d_{gap}=2$ mm, respectively.

A. Gas temperature measurement

One of the most attractive features of nonequilibrium plasmas is that the gas temperature remains close to room temperature (or slightly above room temperature). Here, the gas temperature was determined by analyzing the rotational structure of the N₂ second positive system emission. This rotational structure contains information on the rotational

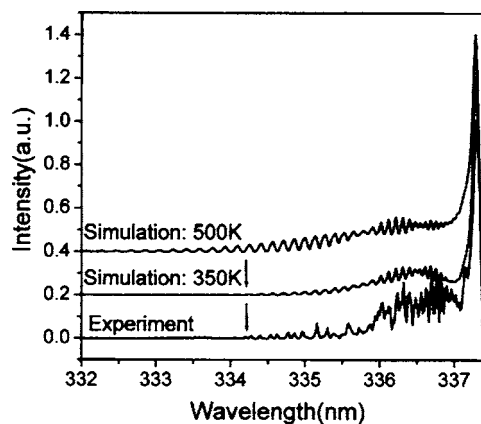


FIG. 2. Experimental and simulation spectra of N₂ second positive 0-0 transition. Applied voltage=9 kV, pulse frequency=1 kHz, pulse width=500 ns, gap distance=2 mm, and operating mixture gas: He(90%) + N₂(10%).

temperature. Because of the low energies needed for rotational excitation and the short transition times, molecules in the rotational states and the neutral gas molecules are in equilibrium. Therefore, the gas temperature can be directly inferred from the rotational temperature.¹⁴ To determine the gas temperature, we compared the experimentally measured spectra with the simulated spectra of the 0-0 band of the second positive system of nitrogen at different temperatures. Figure 2 shows the experimental and simulated spectra results at different gas temperatures for a He(90%) + N₂(10%) operating gas mixture. The spectral resolution is 0.02 nm (grating: 3600 g/mm, slit width: 50 μ m). The curves were intentionally shifted vertically for better separation. Note that the apparent inconsistency of the spectra between 336 and 337 nm is caused by the emission of NO in this band. According to Fig. 2, the higher the gas temperature, the further the band extends towards the short wavelengths. For a gas temperature of 350 K, the simulation results show that the band stops at about 334.2 nm (as shown by the arrow); this is approximately the same as our experimental results. Therefore we concluded that the gas temperature was about 350 K. When the percentages of He (100%–70%) and N₂ (0%–30%) were varied, it was found that the gas temperature did not change appreciably (stayed in the neighborhood of 350 K). In addition, when the pulse frequency was increased from 1 to 10 kHz, the gas temperature only increased relatively slightly (less than 50 K).

B. UV power measurement

UV emission was studied for various mixtures containing different percentages of He and N₂. Figures 3(a)–3(c) show the integrated UV emission power densities for different percentages of He and N₂ and for different pulse frequencies. Figure 3(a) shows that when only helium is used [He(100%) + N₂(0%)] as the operating gas, the emitted UV power density, P_{UV} , increased linearly with the pulse frequency, reaching about 0.18 mW/cm² at 10 kHz. When nitrogen [He(0%) + N₂(100%)] is used as the operating gas,

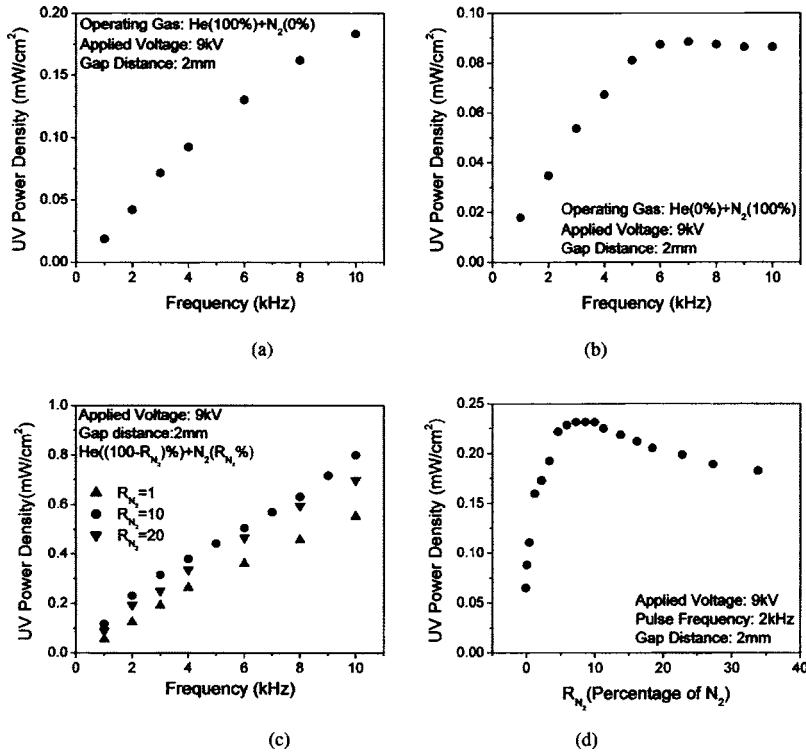


FIG. 3. Integrated UV emission power density vs pulse frequencies for (a) He(100%)+N₂(0%) plasma and (b) He(0%)+N₂(100%) plasma, (c) integrated UV emission power densities for three different percentages (1%, 10%, and 20%) of N₂ in a N₂/He mixture vs pulse frequency, and (d) UV emission power density vs the percentage of N₂ at a fixed pulse frequency of 2 kHz. (R_{N₂}: percentage of N₂)

Fig. 3(b) shows that, in the low-frequency range, P_{UV} initially increased with the pulse frequency, but around 6 kHz, P_{UV} reached a saturation value of about $90 \mu\text{W}/\text{cm}^2$.

To improve the UV emission power density, helium mixed with different percentages of N₂ was then used. Figure 3(c) shows that, for three different percentages of N₂ in a He balance, P_{UV} increases linearly with pulse frequency, and is much higher than that shown in Figs. 3(a) and 3(b). According to Fig. 3(c), P_{UV} is highest when 10% of nitrogen [He(90%)+N₂(10%)] was used. In this case, P_{UV} is more

than four times higher than that of the He(100%) case. To further investigate how the percentage of N₂(R_{N₂}) affects P_{UV} , we measured P_{UV} with He[(100-R_{N₂})+N₂(R_{N₂})] as operating gas mixtures at a fixed pulse frequency of 2 kHz. Figure 3(d) shows that, when the percentage of N₂ was increased from 0% to 5%, P_{UV} increased from about 0.07 to 0.23 mW/cm². Increasing the percentage of N₂ up to about 10% did not greatly affect the value of P_{UV} . However, further increase of N₂ beyond 10% led to a decline in P_{UV} .

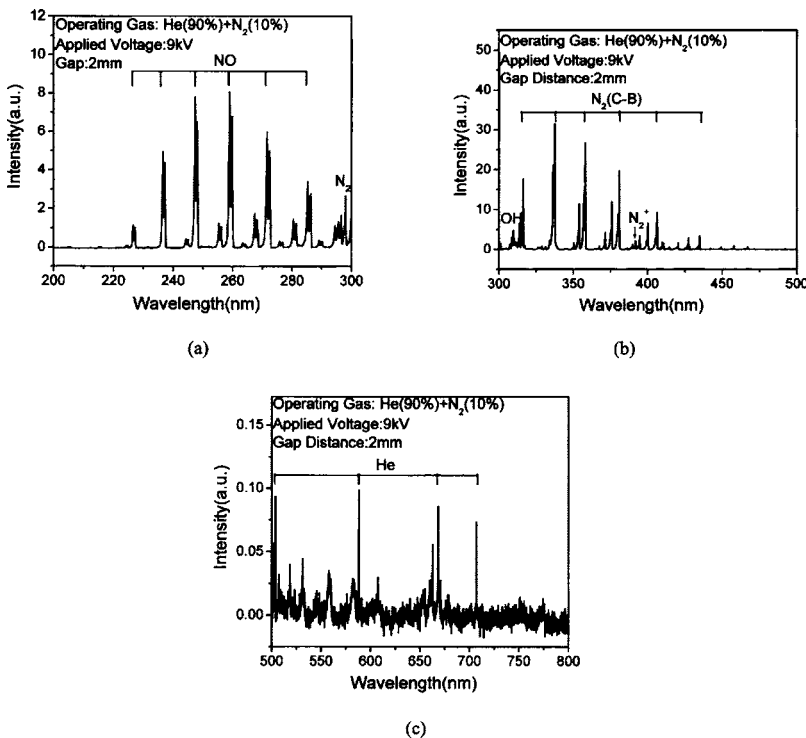


FIG. 4. Emission spectra from (a) 200 to 300 nm, (b) 300 to 500 nm, and (c) 500 to 800 nm for the He(90%)+N₂(10%) plasma. Trace amounts of oxygen are present from residual air.

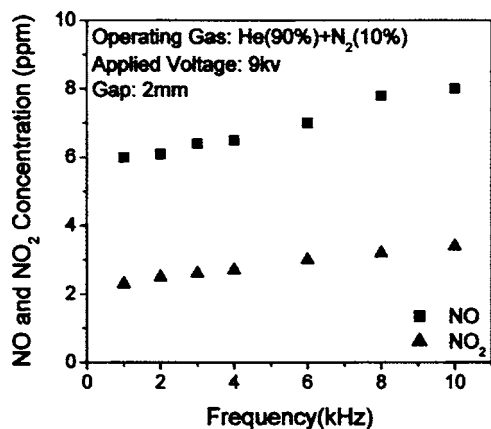


FIG. 5. NO and NO₂ concentrations vs pulse frequency for the He(90%) + N₂(10%) plasma. Oxygen presence is due to trace amounts of residual air.

C. Emission spectra

As shown above, UV emission depends greatly on the gas mixture used in the plasma. To better understand the UV emission characteristics and to identify the presence of chemically reactive excited species in the plasma, emission spectroscopy was used to measure the emission spectra in the 200–800-nm wavelength range (no emission was observed in the 165–200-nm range), under different gas operating conditions. For all of the recorded emission spectra, the pulse repetition rate f was 1 kHz and the operational parameters of the spectrometer were unchanged (grating: 1200 g/mm, slit width: 100 μ m).

Figure 4(a), which shows the emission spectra from 200 to 300 nm for a He(90%) + N₂(10%) mixture, clearly indicates that excited NO radicals are the main source of the UV emission. The N₂(C-B) emission band also contributes to the UV output. Varying the percentages of He and N₂ we observed similar emission spectra but with weaker intensities. Spectra in the wavelength range of 300–800 nm [see Figs. 4(b) and 4(c)] reveal the presence of excited OH, N₂, N₂⁺, and He in the plasma.

D. NO and NO₂ concentrations

Since most of the emission in the UV range was found to originate from NO and since NO is easily oxidized to NO₂ by reaction with O₂, O₃, etc.,^{15,16} we proceeded to measure the absolute concentrations of NO and NO₂. A multigas detector was used for this task. Figure 5 shows the NO and NO₂ concentrations for a He(90%) + N₂(10%) mixture. It shows that both NO and NO₂ concentrations have approximately linear relationships with pulse frequency. The maximum measured NO and NO₂ concentrations were about 8 and 4 ppm, respectively.

IV. DISCUSSION

As stated above, the UV emission from a He(90%) + N₂(10%) plasma is much higher than that when helium or nitrogen alone were used as operating gases. Nitrogen oxide (NO) dominated this emission (where the residual traces of air provided the oxygen to form NO). This improved UV emission can be explained as follows. In order to enhance

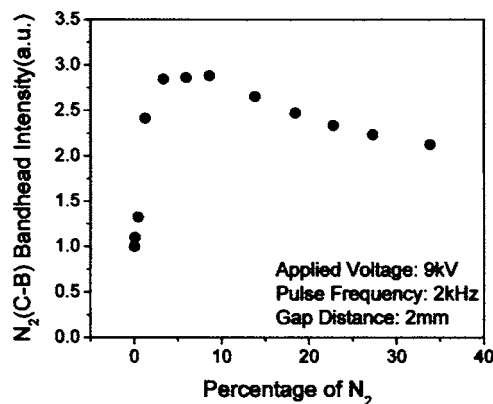


FIG. 6. N₂(C-B) 0-0 transition bandhead (337 nm) intensity vs the percentage of N₂ in a He/N₂ mixture at a fixed pulse frequency of 2 kHz.

UV emission, NO had to be excited to the NO(A² Σ^+) state (excitation energy $E_{NO^*}=5.5$ eV). This can occur via collisions with helium metastables, He*, or with N₂(C³ Π_u) (excitation energy $E_{N_2^*C}=11.1$ eV). According to the spectra that we measured, the N₂(C-B) emission intensity from the plasma with a He(90%) + N₂(10%) mixture was several times higher than that of helium-only or nitrogen-only plasmas. Because the mass of N₂ is seven times higher than that of He, the NO excitation cross section by N₂(C³ Π_u) is likely to be higher than that by He*. Therefore, the plasma with the higher N₂(C³ Π_u) concentration will excite more NO, and as a result emits more UV. To confirm this assumption, we also measured the N₂(C-B)(0-0) transition bandhead intensity (337 nm) for different percentages of N₂ and He mixtures. Figure 6 shows that the emission intensity of this bandhead has a very similar behavior as that of the emitted UV power density shown in Fig. 3(d).

Since most of the UV emission came for the excited NO molecule, the presence of trace amounts of oxygen is therefore required. However, if too much oxygen is added to the gas mixture, the UV emission drops drastically. Figure 7 shows that when just 1% of oxygen was added to a He(89%) + N₂(10%) mixture, the intensity of the NO emission lines between 200 and 300 nm became very weak. Two reasons contribute to this result: (1) In the presence of oxygen NO is quickly oxidized to NO₂. Measurements of the absolute concentration of NO₂ showed a two-order-of-

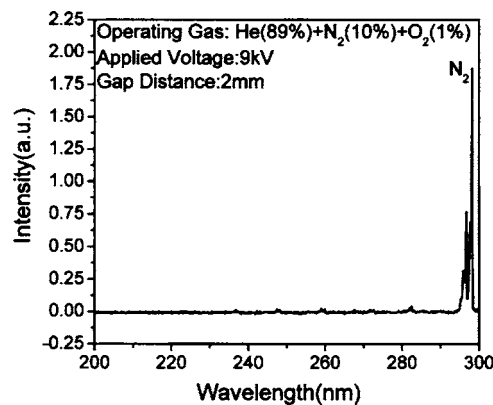


FIG. 7. Emission spectra from a He(89%) + N₂(10%) + O₂(1%) plasma.

magnitude increase (up to 500 ppm of NO₂) when 1% of oxygen was added. (2) Since oxygen has a large electron attachment cross section, it affects the electron energy distribution function in a way that results in less excited species and consequently less UV emission.

Another important observation was that the UV emission power density from a nitrogen plasma was weaker than that from a helium plasma (both in the presence of trace amounts of air). This is because without the presence of He metastables, the only pathway to excite the ground state of N₂ to N₂(C) state is via electron impact.^{17,18} The cross section of this process is smaller than that via He*. So the concentration of N₂(C) state is lower and as a result less NO is excited and less UV is emitted from the nitrogen plasma.

V. CONCLUSION

A specially designed pulsed dielectric barrier discharge was used to generate a diffuse atmospheric pressure plasma. The gas temperature of the plasma was measured to be close to 350 K. The UV emission power density, P_{UV} , measurements show that proper mixture of He and N₂ improves the UV emission dramatically. With 5% to 10% of N₂ mixed with He, P_{UV} reached about 0.8 mW/cm² (for $f=10$ kHz), and was more than four to nine times higher than that for He or N₂ plasmas, respectively. The emission spectra show that most of the UV was emitted by excited NO and N₂. The

emission spectra from a He(90%)+N₂(10%) plasma show relatively high concentrations of excited NO and N₂. In addition to NO, NO₂, and excited OH, N₂⁺ was detected.

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