

대기오염 가스 제거효율 향상을 위한 저온 플라즈마 응용기구 연구

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Non Thermal Plasma Applicable Mechanisms for the Improvement of Air Pollutants Removal Efficiency

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Abstract

A comparative investigation of an experimental and a simulation of chemical kinetics for NO_x removal from dielectric barrier discharges is presented. Several types of dielectric barrier discharges were implemented depending upon the configuration of electrodes. The simulation was based on an approximate mathematical model for plasma cleaning of waste gas. The influence of non uniform distributions of species due to the production of primary active particles in the streamer channel was taken into account. A comparison of observed experimental to the calculated removal efficiency of NO_x showed acceptable agreement.

Key Words : NO_x, non thermal plasma, chemical kinetics simulation, perimental

1. Introduction

One of the promising technologies for destruction or removal of nitrogen oxides (NO_x) from flue gas streams is the non-thermal plasma discharge. Application of the non-thermal plasma for the pollution control is based on a locally generated high electric field. Due to a high electric field, energetic electrons are generated which producing in turn the active components. These components initiate the sequence of chemical reactions involving molecules of the background gases and toxic impurities. These generated radicals reduce and/or oxidize NO_x molecules. The efficiency of NO_x removal depends on the mechanism of applying voltage on electrodes. And, electrical discharge techniques can also be implemented in many different ways, depending on the electrode configurations and the applied power supplies [1,2]. In this paper, using a

simulated gas mixture, the performance of a dielectric-barrier discharge reactor with a coaxial cylinder electrode has been investigated in dry conditions. And, a computer simulation results that based on the experimental data are also presented. The employed simulation modelling of chemical kinetics showed generally in a good agreement with the obtained experimental data.

2. Experimental

In this work, two different types of plasma reactors were implemented for the investigation. The first type reactor with a 20 mm and the second type reactor with a 35 mm outer diameter had a cylindrical glass chamber as a dielectric barrier between the high voltage and ground electrodes. The glass cylinder provides a dielectric barrier to the development of a discharge, so that high voltage

can be applied without breakdown. The external surface of cylindrical reactor glass was covered with an aluminium foil, which forms the ground electrode. High voltage pulses of positive polarity were applied to the inner electrode. The inner electrode of the first type reactor is consisted of a centrally suspended straight wire as shown in Fig.1. The second type reactor is comprised of two cylindrical glass tubes arranged so that the gas flow was directed between two tubes as shown Fig. 2(a). Inside of the inner tube was filled with titanium dioxide (TiO_2) pellets and outside of inner tube was wound with stainless wire to form the electrode as shown in Fig 2(b). Reactor sizes are given in millimeters.

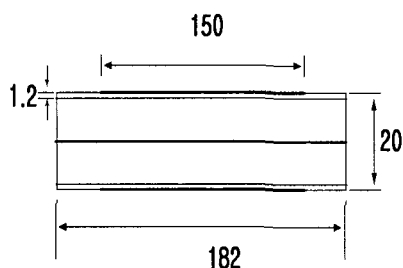


Fig. 1. Schematic diagram of a first type reactor [mm]

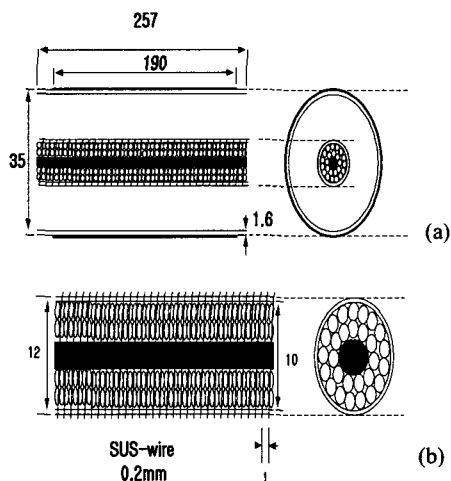


Fig. 2. Schematic diagram of a second type reactor [mm]
(a) geometrical dimension of a reactor (b) inside geometry of inner tube of (a)

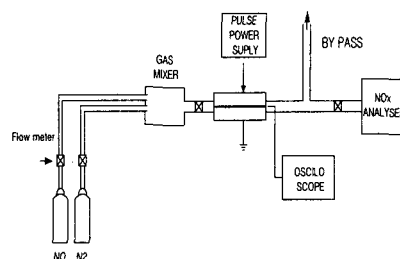


Fig. 3. Schematic diagram of the experimental set-up

A rotating spark gap switch was used to generate the square wave pulses by chopping the DC voltage supplied from DC high voltage generator (Pulse Electronic Engineering, 50 kV). The repetition frequency of pulses was 240 Hz. The output voltage, current and the discharge power were measured using a digital oscilloscope (Tektronix TDS 702A), along with a 1000:1 high voltage divider (Tektronix P6015A) and a current transformer (Tektronix AM503B) with a current probe (Tektronix A6312). A schematic diagram of the experimental apparatus is shown in Fig. 3.

The deposited energy into the gas changed due to applied voltage with an amplitude from 15 to 20 kV at the fixed frequency. The initial NO gas of 1400 ppm with a N_2 balance gas was used and mixed with a dry air to control the variable NO concentration of 300-800 ppm. The flow rate was varied from 2 to 4 l/min. NO and NO_x concentration at the input and the output of reactor were measured with a chemiluminescence $\text{NO}/\text{NO}_2/\text{NO}_x$ analyzer (Advanced Pollution Instrumentation, model 200AU). It is important to point out that in the experiments the gas flowing into the discharge reactor consists of dilute amounts of NO in only N_2 . NO and NO_x concentration were measured in dependence on the deposited energy into the gas.

3. Chemical Kinetics Modelling

For description of experimental results, the approximate mathematical model for plasma cleaning of pollutant gas and the simulation tools of RADICAL had been used [3,4]. The spatial non-uniformity of gas parameters associated with the existence of many

microdischarge channels in a discharge chamber and sequence of discharge pulses were taken into account. The discharge characteristics in experiment such as a specific energy input into a gas per pulse (W_{dc}), a total deposited energy, a number of pulses in the experiment were included in the simulation. Parameters of microdischarge in modelling: a specific energy input into microdischarge channels (W_{st}) and a part of this energy (q) which is spent on the active species production, a fraction of the reactor volume filled by microdischarges ($F_0=W_{dc}/W_{st}$) have been chosen according to the model [3,4]. The parameter q depends on the set-up design and includes some deviations from the theoretical assumptions. It must be adjusted in order to reach agreement with experimental and calculated results for all flow rates for the same experimental set-up and matrix gas composition. In given work the calculations were made for the average electric field of microdischarge $E=100$ kV/cm. This value was chosen because a high voltage was used in these experiments, and, on the other hand, to satisfy a characteristic value of electron density ($\sim 10^{14}$ cm $^{-3}$) and density of filling ($F_0 \sim 10^{-2}$) in a barrier discharge. The concentrations of active species were defined using G-factors corresponding to $E = 100$ kV/cm, and the specific energy for active species production of qW_{st} . G-factor is the number of particles of each sort in dissociative process per 100 eV of absorbed energy. G-factors from literature were properly refined to obtain the G-factors for the considered gas composition.

4. Results and Discussion

It was assumed that only part ($q=0.75$) of specific energy $W_{st} = 0.0075$ J/cm 3 is spent to produce the active species in this barrier discharge. Under these conditions the concentration of nitrogen atoms and electrons were 3×10^{14} cm $^{-3}$ and 3.6×10^{14} cm $^{-3}$ correspondingly. The value of F_0 is changed from 10^{-2} up to 10^{-1} in dependence on the value of W_{dc} for the first and second types of reactors. The value of W_{dc} was changed from 1.7×10^{-4} J/cm 3 up to 8×10^{-4} J/cm 3 . Since the electric field is not known exactly, the sensitivity of its value had been examined. The calculations showed that in general only concentration of

nitrogen atoms (N) is the value, which determines the agreement between calculations and experimental data.

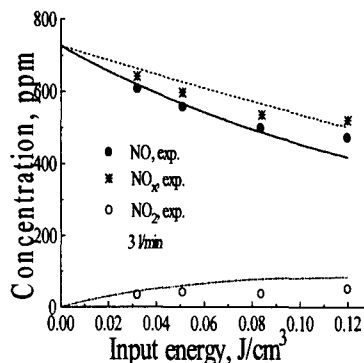


Fig. 4. NO conversion in the discharge chamber for the first type reactor; curves: result of the modelling [3l/min, 730 ppm]

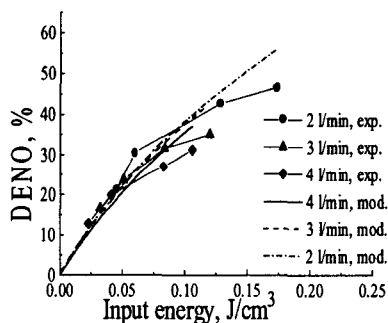


Fig. 5. Conversion of NO for several flow rates and different initial NO concentrations for the first type reactor: 2 l/min, 699ppm; 3 l/min, 730 ppm; 4 l/min, 771 ppm.

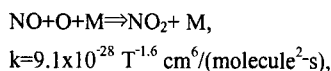
The comparative results of experimental data and calculations in Fig.4 show NO conversion for a flow rate of 3 l/min and initial NO concentration of 730 ppm. Fig.5 illustrates NO conversion ratio for the different initial NO concentrations and flow rates. In general the results of modelling have revealed satisfactory agreement with experiments. Some discrepancy observed at high deposited energy is connected to the heating of gas at 80-100°C (experimental value).

Four different types of discharge depending on method of voltage applying were considered for this reactor. Three different discharges were implemented for electrode configurations: a-apply, b-earth; a-earth, b-apply; a-earth, b-apply, c-earth have been described with identical parameters of microdischarge, where "a" is a metallic rod inside of small tube, "b" is SUS-wire, "c" is a foil. It was assumed that $W_{st} = 0.0075 \text{ J/cm}^3$ and $q=0.25$. In this case, the concentration of nitrogen (N) atoms and electrons were equal to 10^{14} cm^{-3} and $1.2 \times 10^{14} \text{ cm}^{-3}$ correspondingly. The fourth discharge was generated as an electrode configuration of b-apply, c-earth, and the value of q was accepted 0.4 for the same value of W_{st} . The concentration of nitrogen (N) atoms and electrons were $1.6 \times 10^{14} \text{ cm}^{-3}$ and $1.9 \times 10^{14} \text{ cm}^{-3}$.

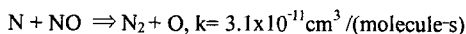
Decomposition efficiency of NO and NO_x are presented for a discharge with electrode configuration of a-earth, b-apply, c-earth in Fig.6 for several flow rates and different initial NO concentrations. In general, taking into account of some possible experimental discrepancies, the modelling is in a good agreement with experiments, especially for NO removal efficiency.

Some difference at the large amount of deposited energy apparently is connected with a local heating of the gas in a microdischarge channel, which is more than the typical average temperature in a gas (final temperature measured in experiments was approximately 380 K).

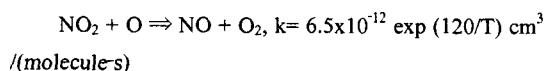
The difference between experimental and calculated results for NO₂ concentration is more appreciable for the discharge configuration of a-earth, b-apply, c-earth than for the configuration of b-apply, c-earth. This may be connected with a non-uniform filling by microdischarge channels of a reactor volume. The value F_0 is larger by a factor of 2-4 for a configuration of a-earth, b-apply, c-earth. In this case the discharge is very strongly located near to the central electrode and more intensive than in a configuration of b-apply, c-earth (observed fact in experimental). In the region of central electrode the temperature is high and the rate constant in reaction



will be smaller, as a consequence of NO₂ concentration will also be rise slower. Whereas the rate constant of conversion NO into N₂ will be same for several types of discharges :



The integral input of the reaction



will be smaller by a factor of 2-4. The rate constants of reactions were utilised from many sources and are quoted in the reference of [3,4]. One can see that the concentrations of NO and NO₂ depend on temperature. Under the large amount deposited energy, the temperature increases, the conversion of NO to NO₂ decreases, and the

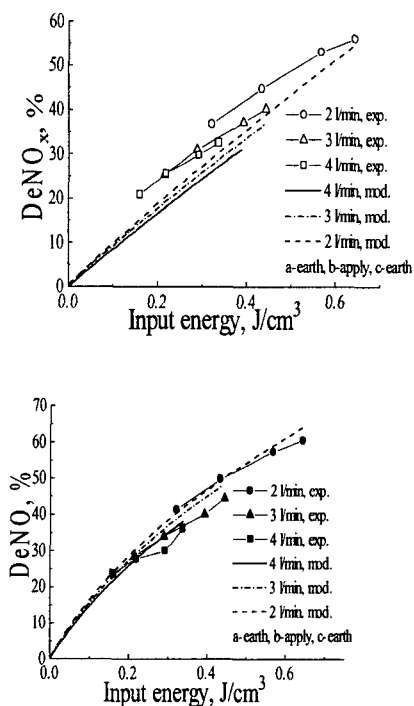


Fig. 6. Decomposition of NO_x and NO for a discharge configuration of a-earth, b-apply, c-earth for several flow rates and different initial NO concentrations for the second type of reactor: 2 l/min, 701 ppm; 3 l/min, 738 ppm; 4 l/min, 781 ppm.

concentration NO_2 almost does not change. The main conversion of NO in a mixture $\text{N}_2\text{-NO}$ is conversion to N_2 and O_2 .

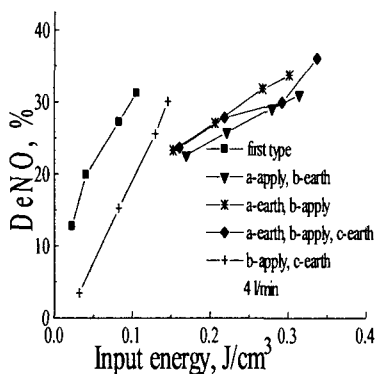


Fig. 7. NO decomposition efficiency for discharge configurations. $[\text{NO}]_0=780$ ppm smaller than that for reactors in [5].

Fig. 7 presents the comparison of NO and NO_x decomposition efficiency for several types of discharge in the first and second types of reactors for the flow rate of 4 l/min and similar initial NO concentration. The similar results were obtained for other flow rates. One can see that a configuration of b-apply, c-earth is more effective than other discharge configurations for a second reactor. Their efficiency is approximately similar and equal to 230 eV for removal of one NO molecule. For configuration of b-apply, c-earth the average energy cost is near to 170 eV. This value for the first type reactor was equal to near 100 eV in $\text{N}_2\text{-NO}$ mixture and appreciable smaller than that for reaction in [5].

5. Conclusions

Removal efficiency of NO_x in N_2 was investigated for the plasma reactor when a high pulse voltage could be applied without breakdown. The results of modelling are close to experimental results. The NO removal depended

on initial NO concentration, flow rate, deposited energy, electrode configurations of discharge and temperature. The better reduction NO and NO_x performance were obtained in lower initial NO concentrations and decreasing gas flow rate. The discharge with straight wire configuration of inner electrode was more effective than other types of discharges. For real mixtures (air, flue gas and others) this reactor may be very effective because the energy cost for N_2 dissociation is much lower than that in air or flue gas for O_2 or H_2O for the same electron mean energy. In correspondence with experiment, such type of reactor could be more suited at the lower input energy when an increasing temperature is small. A good agreement between experimental data and calculations gives opportunity to simulate the removal process in these types of reactors and predict of its efficiency for other gases.

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