DEGRADATION OF REACTIVE ORANGE 16 USING A PROTOTYPE ATMOSPHERIC-PRESSURE NON-THERMAL PLASMA REACTOR†

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Abstract. A prototype atmospheric pressure non-thermal corona plasma reactor system is developed and tested for the removal of commercial textile reactive dye from water. The dye can be completely degraded in water by the presented reactor system in the initial concentration range of 10-100 mg dm\(^{-3}\). Dye degradation rate decreases with the increase of initial dye concentration and pulse frequency. The pH of treated solutions decreases with the increase of treatment time and with the decrease of the applied frequency. Solutions electrical conductivity increases with an increase of treatment time and with the decrease of the applied frequency. The decolorization reactions follow the pseudo-first kinetics order. The changes in the treated dye solutions compositions are the consequences of injection of plasma generated reactive species from gas into the liquid through the interfacial zone.

Key words: plasma, air, dye, degradation, pH, conductivity

1. INTRODUCTION

The interaction of non-equilibrium plasmas with a liquid state is applied in many areas including environmental applications, especially in the wastewater treatments (Bruggeman and Leys, 2009).

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Synthetic organic dyes are very stable and resistant to light, heat, common oxidizing agents and aerobic biological degradation, which makes them very difficult to remove from wastewater by the conventional treatments (e.g. Sun and Yang, 2003). The Advanced Oxidation Processes (AOPs) are, however, capable to degrade dyes molecules by oxidation via highly reactive, short-living oxygen species with very high oxidative potential, like hydroxyl radical, $•$OH ($E^0 = 2.80$ V).

These processes, in which $•$OH radicals are generated, involve electrochemical oxidation (e.g. Martínez-Huitle and Brillas, 2009), UV radiation in the presence of O$_3$, H$_2$O$_2$, photocatalyst, etc. (e.g. Simonenko et al., 2015) and various types of plasmas, such as pulsed corona discharge, dielectric-barrier discharge, atmospheric non-thermal plasma torch and jet, as well as atmospheric pressure non-thermal plasma corona (Magureanu et al., 2008; Tichonovasa et al., 2013; Bansode, et al, 2017; Chandana et al., 2015).

The plasma generated at air-water interface is a source of strong oxidants, like reactive ion and radical species, electrons, UV radiation, electric field etc. Some of them, such as aforementioned $•$OH radical, are capable of degrading various organics in a water medium.

In this paper, a prototype atmospheric pressure non-thermal corona plasma reactor system and its ability to remove a commercial textile dye Reactive Orange 16 (RO 16) from the water solution is described. The discharge was created in the gas phase using multipin electrode over a liquid, which was used as an anode. The liquid itself was a cathode.

Corona occurred in the strong non-uniformed electric field, on the spikes (conductors with a small radius of curvature), and is characterized with low current intensity and a correspondingly small level of emitted light (e.g. Raizer, 1991).

2. EXPERIMENTAL

2.1. Equipment

A multipin-water electrode geometry was used to generate corona in the air at atmospheric pressure. The experimental set-up (Figure 1) consists of a plasma DC pulse generator and reactor chamber which were developed and tested in our Research Laboratory for Gas Discharges. The corona discharge was created near the multipoint anode, composed of stainless steel needles in the gas phase above the liquid. The electrode contains 97 needles and their mutual distance was 2.5 mm. They were soldered on the circular printed circuit board (PCB), with a diameter of 29 mm, all at the same potential. The surface of the conductive liquid acts as a cathode for this discharge and surface area of the liquid electrode is 16.61 cm$^2$. The reactor cell is made of polyvinyl chloride (PVC) in a cylindrical shape. The HV ratios and the frequency (in the range from 10 to 100 kHz) of the pulses were controlled. The distance (d) from the anode to the liquid surface was 16 mm in these experiments. The electrical current of discharge was measured by the multimeter Iskra Unimer 06. The discharge current density was calculated based on the measured discharge current values and the surface of the liquid electrode, and it was 9.15 $\mu$A/cm$^2$. The volume of the samples was 10 cm$^3$. 
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Fig. 1. Schematic of the experimental set-up

2.2. Materials and Methods

RO 16 was obtained from Farbotex (Italy) and used without further purification. All solutions were prepared in deionized water (18.2 MΩ, Smart2Pure, Thermo Scientific, USA).

| Table 1. The main characteristics of RO 16 |
|-------------------------------------------|
| Dye                                      | Chemical structure | Molar mass (g mol⁻¹) | Λ_max (nm) |
|-------------------------------------------|--------------------|----------------------|------------|
| Reactive Orange 16                        |                    | 617.54               | 493        |

2.3. Experimental Procedure

Dye decolorization experiments were carried out at an ambient temperature of 20±0.5 °C (which was held constant during the experiments, at various initial dye concentrations. RO 16 solutions of 10.0, 20.0, 50.0, 80.0 and 100.0 mg dm⁻³ were prepared separately by dissolving 1000.0 mg of a powdered dye in 1 dm³ of water (stock solution) and further diluting required amounts of stock solution to obtain the required concentration dye solutions. The samples of the solutions were taken at the certain time intervals and measured by UV-VIS spectrophotometer Shimadzu UV-1650 PC (Shimadzu, Japan). The percentage of decolorization was calculated using the equation:

\[
\text{Percentage of decolorization (\%)} = \left(1 - \frac{C}{C_0}\right) \times 100 \tag{2.1}
\]

Where \(C_0\) is the initial dye concentration and \(C\) is the dye concentration at the time \(t\). All the experiments were done starting at the solutions initial native pH and conductivity.
Electrical conductivity and pH of the solutions were measured during the treatment using pH meter (SensIon3, Hach, USA) and conductometer (SensIon5). All experiments were conducted in triplicate and gave reproducibility of within 2%.

3. RESULTS AND DISCUSSION

3.1. Plasma chemistry

Fast electrons produced in the non-thermal plasma can have energies of the order 10 eV. The primary source of all reactions is electron impact on the bulk gas molecules. It produces many reactive species as radicals: •OH, •O, and •NO; excited N₂ molecules; and atomic and molecular ions (e.g. O⁺, O₂⁺), which can react further with more stable species (e.g. Nijdam et al., 2012).

Producing of free oxygen radicals by inelastic electron impact on molecules in air:

\[
\begin{align*}
O₂ + e &\rightarrow O⁺ + •O + 2e \\
O₂ + e &\rightarrow O + •O + e \\
O₂ + e &\rightarrow O⁺ + •O
\end{align*}
\]  (3.1)

After that the ozone is created by

\[
•O + O₂ + M \rightarrow O₃ + M, M = O₂, N₂ or Ar.
\]  (3.4)

Formation of NOₓ in the air can be presented in the following way (e.g. Lewis et al., 2003; e.g. Bruggeman et al., 2016):

\[
\begin{align*}
N + O₂ &\rightarrow NO + •O \\
•O + N₂ &\rightarrow NO + N \\
N₂ + O₂ + e &\rightarrow 2NO + e
\end{align*}
\]  (3.5)

The N₂ molecule can be dissociated by electron impact:

\[
N₂ + e \rightarrow N + N + e
\]  (3.8)

Furthermore, the produced NO can react with O₂ (e.g. Bruggeman et al., 2008); e.g. Bruggeman et al., 2008a)

\[
\begin{align*}
•O + NO + M &\rightarrow NO₂ + M \\
2NO + O₂ &\rightarrow 2NO₂ \\
2NO + O₂ &\rightarrow NO₂ + N + O₂
\end{align*}
\]  (3.9 to 3.11)

The removal of NO from the air at low NO concentrations is represented by the Eq. (3.8) followed with the reaction (3.12) (e.g. Bruggeman et al., 2007)

\[
N + NO \rightarrow N₂ + •O
\]  (3.12)

A second important radical is •OH. It is produced in moist gases (e.g., moist air) by the following reaction (e.g. Bruggeman et al., 2008b)

\[
H₂O + e \rightarrow H + •OH + e
\]  (3.13)

Formation of acids HNO₂, HNO₃ is represented by the following reactions (e.g. Bruggeman et al., 2016):

\[
\begin{align*}
NO + •OH &\rightarrow HNO₂ \\
NO₂ + •OH &\rightarrow HNO₃
\end{align*}
\]  (3.14 to 3.15)
In the plasma-liquid interactions, chemical reactions most likely occur in the bulk liquid and the gas phase. At the gas-liquid interface, they appear simultaneously with transport processes.

The reaction

$$2\cdot\text{OH} \rightarrow \text{H}_2\text{O}_2$$

represents the formation of hydrogen peroxide by hydroxyl radical recombination. It can appear in the gas (g), the liquid (aq) or at the gas-liquid interface.

**Fig. 2.** Schematic diagram of plasma chemical processes. Positive ions and UV photons are produced in the discharge zone (D); I - interfacial zone; BL - bulk liquid. Circulations in air and liquid are indicated by multiple arrows.

Corona discharge over the water surface, with a gas gap containing air, can also produce nitrates and nitrites. NO and NO$_2$ are formed in the gas phase from the nitrogen in the air. These products further react with water, hydroxyl radicals and other oxidants in the liquid phase. That way, nitrate, and nitrite are formed, which induce pH changes in the liquid phase (e.g. Locke at al. 2006).

Air plasma discharges also emit UV radiation which is an important process of the plasma produced chemistry in liquid. When UV radiation is absorbed in the liquid water with dissolved gases such as O$_2$, the most important reactions are (e.g. Bruggeman and Leys, 2009):
\[ \text{H}_2\text{O}^{(aq)} + h\nu \rightarrow \cdot\text{OH} + \cdot\text{H} \quad (3.17) \]
\[ \text{H}_2\text{O}^{(aq)} + h\nu (\text{<200nm}) \rightarrow e^- + \text{H}_2\text{O}^- \quad (3.18) \]
\[ e^- + \text{H}_2\text{O}^- + \text{H}_2\text{O}^{(aq)} \rightarrow e^{(aq)} + \cdot\text{OH} + \text{H}_2\text{O}^- \quad (3.19) \]
\[ \text{H}_2\text{O}(g) + h\nu \rightarrow e^- + \text{H}_2 + \cdot\text{O}(1\text{D}) \quad (3.20) \]
\[ \text{O}_3 + h\nu \rightarrow 2\cdot\text{O} \quad (3.21) \]

As can be seen, plasma-induced UV radiation also contributes to a formation of \( \cdot\text{OH} \) radicals.

### 3.2. Effect of Initial Dye Concentration and Decolorization Kinetics

RB 19 shows high resistance to UV radiation alone and \( \text{H}_2\text{O}_2 \) in dark and light (e.g. Mitrović et al., 2012). When RO 16 water solution is exposed to air plasma, being practically a liquid cathode, a fast decolorization is observed. Dye was completely decolorized, in the tested range of initial concentrations. Dye removal efficiency decreases with the increase of its initial concentration, probably because of the higher number or reactant molecules. After the first ten minutes of the treatment, the decolorization efficiency, expressed as a \% of the color removed, was 100\%, 96.76\%, 94.37\%, 91.52\%, and 84.81\%, for the solutions of 10.0, 20.0, 50.0, 80.0 and 100.0 mg dm\(^{-3}\) initial dye concentration, respectively. Total decolorization time varies from about 10 minutes for the lower concentrations to little less than 25 minutes for the highest initial concentration.

Plasma decolorization of RO 16 approximately follows the pseudo-first-order kinetic model, which is described by the equation:

\[ \ln\left( \frac{C}{C_0} \right) = -kt \quad (3.28) \]

where \( C_0 \) is initial dye concentration, \( c \) is a dye concentration after reaction time \( t \), and \( k \) is the reaction rate constant (min\(^{-1}\)). The plot of \( \ln(C/C_0) \) against time scale for the plasma decolorization of RO 16 for a series of the initial dye concentrations ranging from 10 to 100 mg dm\(^{-3}\) (Figure 3) approximately shows a linear decrease with increasing reaction time.

![Fig. 3. Pseudo-first order kinetics model for plasma decolorization of RO 16, \( C_0 \) is given in mg dm\(^{-3}\)](image)
This is confirmed by the relatively high coefficient of determination \( (R^2) \) values of about 0.98-0.99 in the whole range of tested concentrations (Table 2) and is in accordance with the literature, which reports the pseudo-first-order kinetics reaction of organics by various AOPs, including certain types of plasmas (e.g. Chandana et al., 2015; e.g. Chandana and Subrahmanyam, 2016; e.g. Jiang et al., 2012; e.g. Petrović et al., 2015; e.g. Simonenko et al., 2015). These findings strongly indicate that the dye was degraded by the action of highly reactive oxygen species originating from the air plasma, generated in gas and gas-water interface and dissolved in water, eventually leading to a generation of \( \cdot \text{OH} \) radicals, which oxidized and degraded dye molecule through mechanisms and reactions given above and elsewhere in literature (e.g. Bruggeman and Leys, 2009; e.g. Bruggeman et al., 2016; e.g. Barthakur, 1990)

Reaction rate constants decrease with the increase of initial dye concentration (Table 2), falling from 0.421 for the lowest to 0.23 for the highest initial concentration, most likely because the ratio between the amount of “in situ” produced \( \cdot \text{OH} \) radicals and the remaining dye molecules becomes less favorable at the highest dyes concentration.

**Table 2. Reaction rate constants and \( R^2 \) values for decolorization of RO 16**

| \( C_0 \) (mg dm\(^{-3}\)) | Reaction rate constant (min\(^{-1}\)) | \( R^2 \) |
|--------------------------|-------------------------------------|----------|
| 10                       | 0.421                               | 0.9914   |
| 20                       | 0.351                               | 0.9906   |
| 50                       | 0.338                               | 0.9823   |
| 80                       | 0.302                               | 0.9830   |
| 100                      | 0.230                               | 0.9825   |

3.2.1. The pH and conductivity of the plasma-treated dye solutions

Solutions pH decreases during the treatment, starting from its beginning (native) value (which varies from 5.27 to 5.01 for the initial dye concentration of 10 to 100 mg dm\(^{-3}\), respectively) in a very similar manner for all the initial dye concentrations (Figure 4), no matter what their initial pH was. After 10 minutes of treatment, pH falls to about 2.9 for all the treated solution. A decrease of pH was also observed in other plasma treatments of organics in the aqueous medium (e.g. Chandana et al., 2015). Decrease of pH is explained by excitation of \( \text{O}_2 \) and \( \text{N}_2 \) molecules in air, their transformation to \( \text{NO} \) and \( \text{NO}_2 \) molecules via the mechanisms described in the section 3.1 and their further dissolution in water phase, leading to a generation of \( \text{HNO}_2 \) and \( \text{HNO}_3 \), which acidify the solutions (e.g. Chandana et al., 2015; e.g. Shang et al., 2017; e.g. Hentita et al., 2014).

An increase of the solution electrical conductivity (\( \kappa \)) with the increase of treatment time is also observed (Figure 5). Electrical conductivity increases from 11.1 to 77.4 \( \mu \text{S cm}^{-1} \) for the initial dye concentration of 10 to 100 mg dm\(^{-3}\), respectively and it originates from the presence of the dye molecules in water. As the treatment proceeds, conductivity increases for all the treated dye solutions, always being little higher at a certain treatment time for the solutions with the higher initial dye concentration. For the lower initial concentrations, \( \kappa \) remains very similar during all the treatment for the same treatment time, but their corresponding \( \kappa \) values are significantly different from those of the higher initial concentrations solutions at a certain time of treatment because the organic molecules contribute to the overall conductivity of the solutions.
Fig. 4. Changes of pH during the plasma treatment of RO 16

Fig. 5. Changes of electrical conductivity during the plasma treatment of RO 16

However, $\kappa$ increases in the same manner for all the dye solutions and the reason is generally the same as that for the decrease of pH: generation of nitrates and nitrites in dye water solution during the treatment. A very similar result regarding the increase of conductivity was obtained for the treatment of Crystal Violet dissolved in distilled water by DBD induced non-thermal plasma (e.g. Chen et al., 2017) and for the degradation of Methylene Blue dissolved in deionized water by the microwave atmospheric pressure plasma jet (e.g. García et al., 2017). Thus, during the plasma treatment of water medium, if $\text{N}_2$ is present in the gas phase, nitrogen radicals will also be generated and corresponding nitrogen species will be dissolved in the
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water phase, along with the oxygen radicals. The concentration of both kinds increases during the treatment time.

3.3 Effect of frequency on the decolorization rate, pH and electrical conductivity

The 50 mg dm$^{-3}$ dye water solutions were treated at various pulse frequencies ($\nu$) during the same time. The solutions were treated in the frequency range in which is possible to obtain a stable corona; the lowest frequency is the one where corona is still stable, just before it becomes unstable, creating sparks and conductive “plasma channels” in gas phase and the highest frequency is the one when corona becomes the weakest and disappears. As shown in Figure 6, dye decolorization efficiency decreases with the increase of $\nu$, from 68% at 10 kHz to only 26% at 50 kHz for the treatment time of 5 minutes.

![Fig. 6. Effect of applied frequency on dye degradation ($C_0$ (RO 16) =50 mg dm$^{-3}$, t = 5 min).](image)

As discussed before, dye degradation is directly connected with the production of •OH radicals, which indicates that •OH generation in liquid is more intense at lower frequencies, probably because at the higher frequencies the pulses are not long enough to accelerate the generated electrons which strike the air constituents, creating radicals; at lower frequencies, the regime is closer to DC, which means that electrons have more time between impulses to accelerate and strike with higher kinetic energy, or have more successive collisions. The pH of the treated solutions is the lowest at 10 kHz (3.07) and it starts to significantly increase with the increase of frequency, reaching its maximum of 3.72 at 50 kHz. Having in mind that the pH reflects the concentration of HNO$_2$ and HNO$_3$ in solutions, which originate from the plasma nitrogen radicals, the results indicate that the production of nitrogen radicals becomes less intense as the treatment frequency increases, probably due to the same reason as for the oxygen radicals, including longer pauses between the pulses on lower frequencies when recombinations and forming of NO and NO$_2$ may be more probable.

Solution conductivity decreases with the increase of treatment frequency (Figure 7) basically reflecting the same phenomenon as pH, confirming the decrease of generation of nitrogen species in liquid with an increase in frequency. Overall, it can be assumed that
with the increase of treatment frequency the generation of both •OH radicals and nitrogen species in the liquid decreases.

![Image](Fig. 7) Effect of applied frequency on solutions pH and electrical conductivity ($C_0$ (RO 16) = 50 mg dm$^{-3}$, t = 5 min).

4. CONCLUSIONS

A prototype atmospheric pressure non-thermal corona plasma reactor system was successfully applied for the degradation of a commercial reactive textile dye from water. The dye was completely degraded in the tested concentrations range. Dye removal efficiency increases with the increase of treatment time and its removal rate decreases with the increase of initial dye concentration. It proceeds via oxidation of plasma-generated active oxygen species dissolved in liquid, following the pseudo-first kinetics order. Dye solutions pH decreases and their conductivity increases with the increase of reaction time due to the generation of HNO$_2$ and HNO$_3$ caused by dissolution of plasma-generated active nitrogen species. Dye degradation efficiency and solutions electrical conductivity decreases and the pH increases with the increase of treatment frequency, meaning that generation of both active oxygen and nitrogen species in liquid decrease when the treatment frequency increases.

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DEGRADACIJA REAKTIVNE NARANDŽASTE BOJE RO16 KORIŠĆENJEM PROTOTIPA KORONA REAKTORA SA HLADNOM PLAZMOM NA ATMOSFERSKOM PRITISKU

Prototip atmosferskog korona reaktora sa hladnom plazom razvijen je i testiran za razgradnju komercijalne reaktivne tekstilne boje u vodi. Boja može biti potpuno razgrađena primenom navedenog plazma reaktora u opsegu početnih koncentracija od 10-100 mg dm⁻³. Brzina razgradnje boje opada sa porastom početne koncentracije i povećanjem primenjene frekvencije impulsa. Vrednost pH tretiranih rastvora opada sa porastom vremena tretmana i smanjenjem primenjene frekvencije. Električna provodljivost raste sa porastom vremena tretmana i sa smanjenjem primenjene frekvencije. Reakcije obezbojavanja slede kinetiku pseudo-prvog reda. Promene u sastavu tretiranih rastvora su posledica prelaska plazma generisanih reaktivnih vrsta iz gasa u tećnost kroz grančani sloj.

Ključne reči: plazma, vazduh, boja, degradacija, pH, provodljivost