Generation of reactive species in water film dielectric barrier discharges sustained in argon, helium, air, oxygen and nitrogen

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Abstract
Activation of liquids with atmospheric pressure plasmas is being investigated for environmental and biomedical applications. When activating the liquid using gas plasma produced species (as opposed to plasmas sustained in the liquid), a rate limiting step is transport of these species into the liquid. To first order, the efficiency of activating the liquid is improved by increasing the ratio of the surface area of the water in contact with the plasma compared to its volume—often called the surface-to-volume ratio (SVR). Maximizing the SVR then motivates the plasma treatment of thin films of liquids. In this paper, results are discussed from a computational investigation using a global model of atmospheric pressure plasma treatment of thin water films by a dielectric barrier discharge (DBD) sustained in different gases (Ar, He, air, N2, O2). The densities of reactive species in the plasma activated water (PAW) are evaluated. The residence time of the water in contact with the plasma is increased by recirculating the PAW in plasma reactor. Longer lived species such as H2O2aq and NO3−aq accumulate over time (aq denotes an aqueous species). DBDs sustained in Ar and He are the most efficient at producing H2O2aq, DBDs sustained in argon produces the largest density of NO3−aq with the lowest pH, and discharges sustained in O2 and air produce the highest densities of O3aq. Comparisons to experiments by others show agreement in the trends in densities in PAW including O3aq, OHaq, H2O2aq and NO3−aq, and highlight the importance of controlling desolvation of species from the activated water.

Supplementary material for this article is available online

Keywords: plasma activated water, dielectric barrier discharge, modeling plasma liquid interactions

(Some figures may appear in colour only in the online journal)
1. Introduction

Low-temperature atmospheric-pressure plasmas (APPs) are being investigated in applications from material processing to biomedicine and agriculture. The common interest in many of these applications is the interaction of APPs with liquids to activate or produce reactive species in the liquid. Plasma activation of liquid can be facilitated with the plasma in or in contact with the liquid which generally falls into three categories: (1) discharges sustained inside the liquid [1, 2], (2) discharges in the gas phase inside the liquid, such as plasmas in bubbles [3, 4], or (3) plasmas in gas in contact with liquid [5]. In the last approach, dielectric barrier discharges (DBDs), jets, corona, or gliding arcs have been investigated as a means to transfer plasma activated species into the liquid, which affects not only their chemical reactivity but also their conductivity and acidity.

The transport of gas phase species into the liquid is optimized by maximizing the surface area in contact with the plasma. With the goal of activating the volume of the liquid, the process is typically optimized by increasing the surface-to-volume ratios (SVR) of the surface receiving plasma activated species and the volume of liquid being activated. A reactor with a thin water film takes advantage of a high SVR and is therefore frequently studied in water decontamination applications. The interface is the most activated layer where the decomposition of pollutant occurs. In a thin liquid film, the diffusion of pollutant to the activated interface occurs more rapidly due to the larger gradient, while also risking desolvation of those species as well.

In plasmas sustained in air, or having oxygen and nitrogen impurities, reactive oxygen species (ROS) and reactive nitrogen species (RNS) that are generated in the plasma transport to the surface of and solvate into the liquid. Although short lived aqueous species such as hydroxyl (OHaq) are important in many applications, aqueous species such as hydrogen peroxide (H2O2aq), nitrite (NO2aq−), nitrate (NO3aq−) and ozone (O3aq) are often the end products of plasma activation of water due to their longer lifetimes and are effective in antibacterial applications [6–8] (the subscript aq denotes an aqueous species).

Plasma based water decontamination has been investigated for a wide range of contaminants including dyes and organic compounds such as methylene blue and phenol [9, 10]. Malik [11] described different plasma reactors and their energy efficiency for water purification applications, and suggested that pulse powered reactors operating on liquid sprays are the most energy efficient for producing plasma activated water (PAW). Foster [12] reviewed the progresses and needs in plasma-based water treatment as well as the technical challenges to scale-up and implementation. Plasma remediation of hazardous chemicals in water such as dioxin, PCB, and TNT has been demonstrated. For example, DBDs have been shown to be effective in destruction of dioxin-like compounds—a highly toxic and environmentally persistent pollutant—due to the efficient production of UV radiation, solvated electrons, and OH radicals [13]. OH is a strong oxidant which reacts with most organics and several inorganic compounds and plays an important role in the removal of pollutants in water treatment applications [14].

With the goal of maximizing the efficiency of activating water by increasing the SVR, systems have been investigated where the APP contacts a circulating thin water film, typically covering one of the electrodes. The thin film has a large SVR while circulating the water enables buildup of long lived species in the water. Ognier et al [15] experimentally investigated DBD treatment of a thin water film (0.14 mm thick) to remove acetic acid, phenol and heptanol. Results of a computational fluid dynamic model indicated that the mixing in the liquid film due to interfacial momentum transfer between gas and liquid can accelerate mass transfer of contaminants from the liquid bulk toward the interface and destruction of pollutants [15]. Andre et al investigated a DC discharge in which plasma was ignited in air between two films of flowing tap water [16, 17].

Kuraica et al developed a DBD reactor with a circulating water film flowing over the dielectric surfaces of the coaxial system [18]. The discharge was sustained between the inner dielectric-covered electrode and the water-covered dielectric of the inner electrode operating in air, N2, O2, Ar and He. This falling water DBD (FW-DBD) reactor was used to investigate water purification and removal of pollutants such as phenol [19], arsenic [20], textile dyes [21] and gas phase pollutant such as NOx and SO2 [22]. For example, with a FW-DBD operating in air at 115 W, a 6 min plasma treatment of water resulted in significant removal of 2-chlorophenol (91%), 4-chlorophenol (85%) and 2,6-dichlorophenol (65%) [23]. The ability to customize the type and ratio of reactive species in these systems using different feed gases is attractive to target specific pollutants or destruction of organisms [24]. For example, OH and H2O2 are typically thought of as being the major biocidal components. However, measurement of gas phase and dissolved ozone in an indirect DBD sustained in air indicated a relatively high concentrations of O3 (below about 0.2 W cm−2) and that 5 s of plasma treatment resulted in a completely E. coli bacteria inactivation [25]. The ability to selectively choose the active species would be an advantage.

In this paper, we report on results of a computational investigation using a global model, of a low frequency FW-DBD reactor patterned after the work of Kovačević et al [18] operating at atmospheric pressure in argon, helium, air, oxygen, and nitrogen, with trace air and water impurities. PAW is circulated through the reactor (that is, any volume of water sees several transits through the discharge) to better emulate the experiment and to evaluate ROS and RNS accumulation by increasing the residence time. The gas and liquid phase chemical reaction mechanisms will be discussed, with liquid phase densities compared to experiments for validation.

We found that FW-DBDs sustained in argon are the most efficient gas in the production of hydrogen peroxide in gas and water, generating 0.23 mM of H2O2 over 10 cycles of treating the PAW in agreement with experiments [18]. (M is moles/liter.) Production of gas phase hydroxyl is highest in argon plasma; however, the highest solvated hydroxyl density occurs with air plasmas, while the highest solvated ozone density occurs when operating the FW-DBD in oxygen. Being
short lived, the residual density of OH and OH$_{aq}$ does not increase with re-circulation. The density of HO$_2$ in argon and helium plasmas is several factors higher than in oxygen and air plasma due to the relative abundance of H atoms. Air plasma is the most efficient gas in the production of gas phase NO$_2$ and N$_2$O as well as ONOOH$_{aq}$ in PAW. In all gases, density of NO$_2$$_{aq}$ increases linearly with re-circulation and the highest concentration was found in plasma sustained in argon (1.2 mM) followed by air (0.45 mM). With the presence of RNS, nitrate is the major contributor to the acidity of PAW. Therefore, the lowest pH was obtained in argon followed by air. Results from the model agree systematically and quantitatively with experiments for all species except for O$_{3aq}$. For O$_{3aq}$, agreement is systematic but not quantitative, a result that is attributed to desolvation of O$_{3aq}$ during the afterglow. These results emphasize the importance of controlling desolvation from PAW.

The model used in the investigation is described in section 2 followed by discussion of our computational scaling of FW-DBDs sustained in Ar He, air, O$_2$ and N$_2$ in section 3. The recirculation of PAW is discussed in section 4. Concluding remarks are in section 5.

2. Description of the model and reactor geometry

The model used in this investigation is a 0-dimensional global simulation, GlobalKin, which includes gas flow, transport of plasma activated species to-and-from the plasma-liquid interface and in-liquid reactions. The model is essentially the same as described in detail in [26] and so will be discussed only briefly here. GlobalKin consists of a set of rate equations whose integration provides densities of species in the gas and liquid phases as a function of time or, in plug-flow mode, as a function of position. The average electron energy in the gas phase plasma is obtained from an electron energy conservation equation. Electron impact rate coefficients and transport coefficients are obtained from solutions of Boltzmann’s equation for the electron energy distribution, processed into a periodically updated look-up table as function of electron temperature.

The transport of species between the gas and liquid are limited by Henry’s law equilibrium for the neutral species, while ions and photons from the gas phase can directly enter into the liquid. By definition, at Henry’s law equilibrium the density of a gas dissolved in a liquid is proportional to the density of that species in the gas phase. At equilibrium, the liquid is saturated with the gas phase species. In the model, if the liquid is under-saturated for a particular species based on the Henry’s law constant, transport of that gas phase species is allowed into the liquid. If the liquid is super-saturated (that is, the in-liquid density exceeds that at equilibrium), the net transport of the species is from the liquid to the gas. For instance, species with a small Henry’s law constant saturate quickly in the liquid and its aqueous counterpart has a lower density. Species having a large Henry’s law constant may never be saturated in the liquid, and continue to flow from the gas into the liquid.

Table 1. Henry’s law constants [26].

<table>
<thead>
<tr>
<th>Species</th>
<th>h (unitless)</th>
<th>Note</th>
</tr>
</thead>
<tbody>
<tr>
<td>H$_2$O</td>
<td>$1.92 \times 10^6$</td>
<td>–</td>
</tr>
<tr>
<td>HO$_2$</td>
<td>$1.32 \times 10^5$</td>
<td>–</td>
</tr>
<tr>
<td>OH$_2$</td>
<td>$6.20 \times 10^2$</td>
<td>–</td>
</tr>
<tr>
<td>H$_2$O$_2$</td>
<td>$1.80 \times 10^{-2}$</td>
<td>–</td>
</tr>
<tr>
<td>O, O$_2$(D)</td>
<td>1</td>
<td>–,b</td>
</tr>
<tr>
<td>O$_2$, O$_2$(v), O$_2$(A), O$_2$(A$_2$)</td>
<td>$3.24 \times 10^{-2}$</td>
<td>a</td>
</tr>
<tr>
<td>O$_3$</td>
<td>$3.00 \times 10^{-3}$</td>
<td>–</td>
</tr>
<tr>
<td>N$_2$, N$_2$(A, A$_2$), N$_2$(v), N, N$_2$(D), N$_2$(a, a$_2$)</td>
<td>$1.60 \times 10^{-3}$</td>
<td>a</td>
</tr>
<tr>
<td>N$_2$O</td>
<td>$6.00 \times 10^{-7}$</td>
<td>–</td>
</tr>
<tr>
<td>N$_2$O$_4$</td>
<td>$3.69 \times 10^{-1}$</td>
<td>–</td>
</tr>
<tr>
<td>N$_2$O$_5$</td>
<td>$4.85 \times 10^{-1}$</td>
<td>–</td>
</tr>
<tr>
<td>N$_2$O$_6$</td>
<td>$5.99 \times 10^{-4}$</td>
<td>–</td>
</tr>
<tr>
<td>HO$_2$NO$_2$</td>
<td>$2.99 \times 10^{-2}$</td>
<td>–</td>
</tr>
<tr>
<td>NO$_2$</td>
<td>$4.40 \times 10^{-2}$</td>
<td>–</td>
</tr>
<tr>
<td>NO$_3$</td>
<td>$2.80 \times 10^{-3}$</td>
<td>–</td>
</tr>
<tr>
<td>NO$_4$</td>
<td>$4.15 \times 10^{-1}$</td>
<td>–</td>
</tr>
<tr>
<td>HNO$_2$, HNO</td>
<td>$1.15 \times 10^{0}$</td>
<td>a</td>
</tr>
<tr>
<td>HNO$_3$, ONOOH</td>
<td>$4.80 \times 10^{0}$</td>
<td>–</td>
</tr>
<tr>
<td>NH$_3$</td>
<td>$1.47 \times 10^{3}$</td>
<td>–</td>
</tr>
</tbody>
</table>

Value corresponds to the first species in the list. Other species were assumed to have the same Henry’s law constant.

Approximated. Species reacts quickly in water and will not reach Henry’s law saturation.

Approximated by analogy to NH$_3$. liquid phase reactions. The mechanism includes the species associated with He, Ar and humid air plasmas interacting with water. The gas phase reaction mechanism was adapted from Van Gaens et al [27] for argon-humid air mixtures at atmospheric pressure. Extension to He-humid air and interactions with liquid water are based on the mechanism used by Lietz et al [26], Tian et al [28], and Norberg [29]. The mechanism only includes small water cluster ions [O$_3$$^+$(H$_2$O), H$^+$ (H$_2$O)$_2$, and H$_2$O$^+$ (H$_2$O)] in the gas phase. The additional reactions added to the mechanisms are discussed in the supplementary information (available online at stacks.iop.org/JPD/53/435206/mmedia). The interaction of ions and excited states of He with liquid H$_2$O was estimated based on the branching ratios from the gas phase analogues [28, 30]. Henry’s law constants used in this work are in table 1.

A schematic of the FW-DBD reactor used in this investigation, shown in figure 1, based on the experiments of Kovacević et al [18]. In the experiment, water was pumped from a reservoir, flowed along the dielectric covered surface of the inner electrode, returned to the reservoir and then recirculated through the reactor for ten cycles. The FW-DBD is a concentric 1 cm long cylindrical reactor with two dielectric covered electrodes with inner and outer radii of 10 mm and 14.5 mm making for a 4.5 mm discharge gap. The water film on the inner electrode is 0.22 mm thick. The reactor operates at atmospheric pressure and room temperature. Conceptually, the global model is applied to a control volume or plug of liquid film plus the surrounding gas. The control volume resides in the plasma for 1.56 s to match the experiment. Five gas mixtures were examined—Ar, He, air, O$_2$ and N$_2$, and N$_2$O.
Figure 1. Operating schematic and conditions. (a) Schematic of the flowing water DBD reactor. The DBD has an inner radius of 10 mm, outer radius of 14.5 mm and 4.5 mm gap. The water film is 0.22 mm thick. (b) Power profile used in the model. This pulse is repeated at 600 Hz.

with a flow rate of 5 slm. Since the experiment was operated without special vacuum equipment, air and H₂O impurities were added to the gas mixtures in the simulation. Evaporation from the water layer is included in this model and reaches to a steady state of 2.5% humidity for all gases. The liquid is pure water with 8.93 ppm (based on molecular densities) dissolved N₂ and 4.81 ppm dissolved O₂, the equilibrium values with air. The diffusion length in the gas phase is \( d/\pi = 1.36 \text{ mm} \), where \( d \) is the width of the discharge gap. When including the saturated water vapor, the air gas mixture is \( \text{N}_2/\text{O}_2/\text{H}_2\text{O} = 77.5/20/2.5 \), and helium and argon mixtures are 96.5% pure with 3.5% added humid air impurities (\( \text{N}_2/\text{O}_2/\text{H}_2\text{O} = 0.8/0.2/2.5 \)). Accordingly, the mole fractions of N₂ in the nitrogen plasma is 97.3% and O₂ in oxygen plasma is 96.7%. CO₂ as an air impurity was not included in the reaction mechanism.

In this model, we specified power deposition as a function of time as a 21 W triangular pulse with 40 ns duration at 600 Hz (1.66 ms inter-pulse period). The applied power peaks at the calculated magnitude to provide the measured average power of 21 W to all gases. In the experiment, a sinusoidal voltage at a frequency of 300 Hz was used. Our choice of 600 Hz, 40 ns power pulses acknowledges that experimentally discharge pulses are produced at least twice per cycle, and that the individual discharges are typically only tens of ns long. The experimental dissipated powers were 60, 41, 32, 21 and 16 W for the discharges in air, nitrogen, oxygen, argon and helium respectively, to provide empirically stable operation with the water film and uniformly distribution of microdischarges along the reactor. Therefore, the power in argon matches that of the experiment. Applied power in the model extends for roughly 1500 pulses (2.5 s) after which the system evolves during the afterglow for 10 s (12.5 s total). To simulate PAW recirculation, the final density of gas and liquid species at 12.5 s are used as the initial inputs to simulate the next cycle. The recirculation was conducted 10 times, with each cycle beginning with the final density from the previous cycle at the same operating conditions. By holding the inner and outer electrodes at 300 K, there is no significant gas heating in the base case.

3. Plasma activation of water film

3.1. Plasma properties operating in different gases

The FW-DBD was operated in the 5 gas mixtures (Ar, He, air, O₂, N₂) with an input power of 21 W. The electron density \( (n_e) \) and temperature \( (T_e) \) are shown in figure 2 for the first pulse. A triangular power is applied over 40 ns with the maximum power density of 13.4 kW cm⁻³ at 20 ns. The initial pre-pulse electron density was \( 1 \times 10^8 \text{ cm}^{-3} \). In all gases, the density of electrons increases during the power rise time, peaks shortly after 20 ns at maximum power and decays with a distinctive rate for each gas composition. As the electron density avalanches during the power rise time, the plasma conductivity increases and so the E/N across the plasma decreases which then leads to a reduction in \( T_e \). For instance, in helium \( T_e \) decreases from 8.8 eV to 3.7 eV at the maximum power deposition. The maximum \( n_e \) occurs in helium, \( 3.4 \times 10^{12} \text{ cm}^{-3} \), followed by the argon plasma with \( 9.5 \times 10^{11} \text{ cm}^{-3} \). The lowest \( n_e \) is in nitrogen plasmas, with a peak at \( 1.0 \times 10^{11} \text{ cm}^{-3} \). With power being specified, the electron density is in large part determined by the power dissipation per electron for collisions with the feedstock gases (with impurities) at a given \( T_e \). Power dissipation per electron is lowest in He requiring a higher electron density to dissipate the peak power. Power dissipation per electron is highest in N₂ requiring a lower electron
Figure 2. Electron temperature and density during the discharge pulse for different feed gases. (a) Ar, (b) He, (c) air, (d) O₂, and (e) N₂.

density to dissipate the peak power. Experimental measurements in APPs showed that at a constant applied voltage, current and nₑ are higher in Ar than in He, which agrees with the lowest specific dissipated power in He [31, 32]. Tₑ in all gases sharply decreases during the power fall time so that the power-dissipation per electron decreases.

The decay of electron density shows two distinct rates in these gases, fast (order of tens of ns) and slow decay (up to hundreds of µs). In helium, argon, and nitrogen the electron loss processes are dominated by electron–ion recombination with the rare gas dimer ions and N₂ molecular ions, and dissociative attachment to the water vapor. For air and oxygen
plasmas, attachment to O₂ can be a dominant electron loss process and is largely responsible for the more rapid reduction in electron density for those mixtures. Electron attachment to O₂ in APPs follows two pathways, 2-body dissociative attachment producing O⁻ which has a threshold energy of 3.64 eV and 3-body attachment to form O₂⁻ with thermal electrons. The rate coefficient for 3-body electron attachments to O₂ increases with decreasing Tₑ (<1 eV). Therefore, 3-body attachment is the dominant pathway for electron losses during the power-down portion of the pulse and afterglow when Tₑ is not high enough for a dissociative attachment. For discharge pulses after the first, the buildup of RONS
(reactive oxygen and nitrogen species) such as $O_3$, $N_2O$ and $O_2^-$ (dominated by $O_3(\Delta)$) can introduce additional attachment pathways. $T_e$ remains relatively high (3.1 eV) up to the end of the power pulse in the oxygen and air plasmas, resulting in 2-body attachment being the dominant electron loss process and $n_e$ decreasing rapidly up to the end of the pulse. In argon, helium, and $N_2$ plasmas, $T_e$ decreases during the power fall time to below the threshold for 2-body attachment threshold (down to around 1 eV). With the $O_2$ density being lower and relying on 3-body attachment and dissociative recombination,
the electron decay time increases to hundreds of ns to microseconds.

Densities of short lived ions originating from humid mixtures (with impurities) for Ar, He, air, O₂, and N₂ plasmas for the 1st pulse are shown in figure 3, and for the last, 1500th pulse after a 2.5 s plasma discharge are shown in figure 4. H₂O⁺ is generally the dominant positive ion during the discharge for mixtures other than air and O₂ (that is, for mixtures in which O₂ is an impurity) and O₂⁺ is the dominant ion in the air and O₂ mixtures. These trends result from these ions having the lowest ionization potential and a reasonably large mole fraction of their parent neutral molecule, and so can be produced by rapid charge exchange. The main source of H₂O⁺ at this point is electron impact ionization of H₂O, charge exchange from ions having larger ionization potentials (HeH⁺, He⁺, He₂⁺ in helium; Ar⁺ and Ar₂⁺ in argon, and N₂⁺ in N₂) and Penning ionization from the rare gas excited states. With these processes, H₂O⁺ is the first ion to rise in density in Ar, He and N₂ plasmas. Those mixtures with large mole fractions of O₂ have O₂⁺ as the ion with the initially rising density.

The density of O⁻ is limited to the pulse duration in O₂ and air indicating that the main pathway to its formation is the 2-body dissociative attachment (with a threshold energy of 3.6 eV). The 3-body attachment at low Tₑ in other gases results in later formation of O₂⁻. Formation of O₂⁻ is dominantly through the charge exchange reaction of O₂ or O₁ with O⁻ or through 3-body attachment to O₂:

\[
O^- + O_2 \rightarrow O^-_2 + O,
\]  

\[
e^- + O_2 + M \rightarrow O^-_2 + M,
\]  

\[
O^- + O_3 \rightarrow O^-_5 + O_2.
\]

The maximum density of O₂⁻ occurs in oxygen plasmas in which it is the most abundant ion. In oxygen and air plasmas, O₂⁻ density raises and saturates soon after the power terminates with the dominant pathway being charge exchange reactions (equation (1a)). However, in other gases (Ar, He, and N₂), O₂⁻ peaks hundreds of ns later when Tₑ is below 0.04 eV and so the 3-body reaction is the primary pathway. These trends for the density of O₂⁻ match well with the time history of nₑ and Tₑ for each gas.

Short lived species including ions and vibrationally and electronically excited species, are formed and consumed on a pulse-to-pulse basis by electron impact and are quenched or consumed by reactions prior to the next pulse. O₃⁻ is formed relatively late through the pulsing after the density of O₂ has achieved a reasonably large value. The lowest density of O₃⁻ is in nitrogen and highest density is in oxygen due to higher density of O₂⁻ and O⁻ and larger densities of O₂. The major pathways to O₃⁻ formation are:

\[
O^- + O_2 + M \rightarrow O^-_3 + M,
\]  

\[
O^- + O_3 \rightarrow O^-_5 + O,
\]

\[
O^-_2 + O_3 \rightarrow O^-_5 + O_2.
\]

In oxygen and air plasmas, O₃⁻ decays with two rates. Initially during early pulses, 2-body reactions (equation (2b) and (2c)) contribute to O₃⁻ formation. However, at later times, 3-body processes form O₃⁻ (equation (2a)) and extends its lifetime to hundreds of μs. Although, over 1500 pulses, O₃ accumulates in the plasma, resulting in direct charge exchange from O⁻ and O₂⁻ to O₁ being the dominant source of O₃⁻ in oxygen and air plasmas.

The density of OH⁻ is higher in helium and argon plasmas and is formed at the beginning of the discharge by dissociative attachment to H₂O (e + H₂O → OH⁻ + H). This reaction forming OH⁻ has a high threshold energy (4.5 eV) and so has a higher rate coefficient in Ar and He due to the larger Tₑ during the plasma avalanche. The density of OH⁻ peak at the maximum of the input power and gradually decays, producing the highest density in He discharges. OH⁻ develops later in oxygen and air due to the lower Tₑ, and decays earlier in oxygen (a few μs) due to charge exchange to form O₃⁻.

The densities of O₂⁺, H₂O⁺, and O⁻ generally do not change with successive pulsing while densities and lifetime of ions such as O₂⁻, O⁻, and OH⁻ do change over pulsing. The relatively constant densities of O₂⁺, H₂O⁺, and O⁻ are a consequence of both their production and quenching being from reactions with species that are initially in the gas mixtures. The changes in densities of O₂⁻, O⁻, and OH⁻ suggest the formation and quenching reactions for these ions are sensitive to the accumulation of products on a pulse-to-pulse basis. For instance, in plasmas sustained in helium and argon, over the 1500 pulses the lifetime of O₂⁻ decreases by a factor of about 10 and the density of O₂⁻ increases by a factor of 2 due to the accumulation of O₁ (equations (2b) and (2c)). In air, O₂, and N₂ plasmas, the lifetime of OH⁻ decreases significantly with pulsing, a decrease that is more pronounced in O₂ which can be due to the charge exchange reaction with O₃.

### 3.2. ROS and RNS in the gas phase

The water film thickness is 0.22 mm with 2.5 s residence time per pass through the reactor and total plasma treatment time of 25 s over ten cycles. In this section formation of RONS in gas and water over one cycle (pass) will be discussed. ~2.5 s of plasma exposure followed by a 10 s afterglow. Densities of neutral ROS and RNS in gas phase are shown in figures 5 and 6 respectively for each gas. With the multiple pulses, transient species such as O, OH, HO₂, H, N, and NO are produced and consumed from pulse-to-pulse to generate more stable species that approach a quasi-steady state. The relatively low frequency, 600 Hz, can work in favor of solvation and reactions in liquid phase when the diffusion time of the gas phase species is smaller than the inter-pulse period. If a gas phase species is likely to react with a product species in the gas phase (e.g. OH reacting with OH to form H₂O₂), having a long interpulse period enables the OH to solvate before encountering a newly
Figure 5. Density of gas phase ROS (reactive oxygen species) during the first cycle (2.5 s of plasma exposure and 10 s of afterglow) for different feed gases. (a) Ar, (b) He, (c) air, (d) O$_2$ and (e) N$_2$. (f) ROS for an air feedstock are shown on an expanded time scale during the plasma period.

The generated density of OH in the gas phase produced by the next pulse. Lietz et al [26] reported that the increase in pulse frequency in DBD plasmas decreases the density of most RONS in the liquid due to their likelihood to react with other RONS in the gas phase before solvating. The exceptions were O$_3$aq and HNO$_3$aq whose gas phase precursors are the end product of reactions in the gas phase, and so benefit from the higher repetition rate. The dominant neutral product in helium and argon plasmas is H$_2$, but in air, oxygen, and nitrogen plasmas O$_3$ is the dominant gas phase product. These species also have low Henry’s law constants and accumulate in the gas phase rather than solvating.
With all gas mixtures being humid, decomposition of water generates OH, H and O through electron impact dissociation of water (dominantly by \( e + H_2O \rightarrow O + 2H + e \) and \( e + H_2O \rightarrow OH + H + e \)) or dissociative recombination of \( H_2O^+ \) or \( HO_2^+ \). The larger densities of \( H_2O^+ \) in argon and helium plasmas produce higher densities of OH and O radicals through the dissociative recombination process. OH is produced and consumed on a pulse-to-pulse basis with the loss being dominated by reactions which form \( H_2O_2 \):

\[
OH + OH \rightarrow H_2O_2, \quad (k = 1.7 \times 10^{-11} \text{cm}^3 \text{s}^{-1}), \quad (3a)
\]

\[
H + HO_2 \rightarrow H_2O_2, \quad (k = 3 \times 10^{-11} \text{cm}^3 \text{s}^{-1}), \quad (3b)
\]

\[
HO_2 + HO_2 \rightarrow H_2O_2 + O_2; \quad (k = 2.2 \times 10^{-11} \text{cm}^3 \text{s}^{-1}) \quad E_a = -600 \text{ K} \quad (3c)
\]

(The rate coefficients, \( k \), are for a gas temperature of 300 K with units of cm\(^3\) s\(^{-1}\) (1 cm\(^3\) s\(^{-1}\) = 6.02 \times 10\(^{20}\) M\(^{-1}\) s\(^{-1}\)) unless otherwise noted.) Apart from electron impact dissociation of \( H_2O \), Penning ionization and dissociative charge exchange also contribute to OH formation in ground and excited states in helium and argon plasmas,

\[
Ar^+_2 + H_2O \rightarrow ArH^+ + OH + Ar, \quad (4a)
\]

\[
He^+_2 + H_2O \rightarrow OH(a) + HeH^+ + He, \quad (4b)
\]

\[
He^+ + H_2O \rightarrow H^+ + OH + He. \quad (4c)
\]

Measurements using laser induced fluorescence (LIF) in a helium pulsed filamentary discharge showed that OH is mainly produced by charge transfer reactions to form \( H_2O^+ \) followed by electron–water ion recombination [33]. The high energy of He excited states can produce Penning ionization in \( O_2 \) and \( N_2 \) which results in a higher density of \( O_2^+ \) and \( N_2^+ \). For instance, density of \( O_2^+ \) is one order of magnitude higher in helium than argon plasmas due to these Penning processes,

\[
He^+ + O_2[N_2] \rightarrow He + O_2^+ [N_2^+] + e \quad (5)
\]

The high reactivity and short lifetime of OH limit the diffusion of OH into the water layer. \( H_2O_2 \) is a more stable ROS that also has the highest density in argon plasmas \((4 \times 10^{14} \text{ cm}^{-3})\) due to the higher density of OH and \( H_2O \). \( H_2O_2 \) quickly reaches a steady state (in less than 1 s) as do its reactants (i.e., OH and \( H_2O \)). The densities of \( H_2O_2 \) and OH are lower in air and nitrogen plasmas \(10^{12}-10^{13} \text{ cm}^{-3}\), due to the lower rates of dissociation of \( H_2O \), and lower rates of \( H \) abstraction from \( H_2O \) by O atoms. The initial decrease in \( H_2O_2 \) density on successive pulsing is due to the initial depletion in OH density. In all gases \( H_2O_2 \) is quickly depleted during afterglow, faster than the pump exhaust, due to solvation into the liquid and dissociation reactions such as \( O + H_2O_2 \rightarrow H_2O + O_2 \) with timescales of seconds.

H atoms are the other by-products of \( H_2O \) dissociation, which then react to form \( H_2 (H + H + M \rightarrow H_2 + M) \). Not being very reactive, \( H_2 \) accumulates over the pulse periods while the density of \( H \) quickly reaches a quasi-steady state. The densities of both \( H \) and \( H_2 \) are high in helium and argon plasmas. One of the main reactions to form hydroperoxyl radicals (\( HO_2 \)) is \( H + O_2 + M \rightarrow HO_2 + M \). The highest density of \( HO_2 \) is in argon plasma which correlates well with the higher \( H \) density of \( 7 \times 10^{13} \text{ cm}^{-3} \). In fact, the density of \( HO_2 \) in argon and helium plasmas is 10–100 times higher than in nitrogen and air plasmas due to the abundance of \( H \) atoms. \( HO_2 \) is consumed through \( OH + HO_2 \rightarrow H_2O_2 + O_2 \) \((k = 4.8 \times 10^{-11}, E_a = -250 \text{ K})\) and \( HO_2 + O_2(\Delta g) \rightarrow OH + O_2 + O \quad (k = 1.7 \times 10^{-11}) \). The higher rates of production of \( O_2(\Delta g) \) in \( O_2 \) and air plasmas reduce the density of \( HO_2 \).

The largest production of \( O_3 \) is in air and oxygen plasmas due to the higher \( O \) densities. The major contribution to production of \( O_3 \), the reaction \( O + O_2 + M \rightarrow O_3 + M \), directly depends on production of \( O \) atoms. As shown in figures 5(d) and (c), the highest \( O \) atom density is in oxygen \((5 \times 10^{14} \text{ cm}^{-3})\) and air plasmas \((8 \times 10^{13} \text{ cm}^{-3})\). Spectroscopy of the FW-DBD also indicated strong emission spectra of atomic \( O \) in the oxygen plasma [18].

Electron impact dissociation of \( O_2 \) and dissociative quenching of \( N_2(A) \) by \( O_2 \) are the major pathways to form \( O \) atoms. The higher \( O \) and \( O_2 \) densities in air and oxygen plasmas yield more \( O_3 \) formation. The \( O_3 \) density in the gas phase reaches \( 6 \times 10^{16} \text{ cm}^{-3} \) in the oxygen plasma, which is more than 2 orders of magnitude higher than in helium and argon. The density of \( O_3 \) accumulates over pulsing in \( O_2, N_2 \), and air plasmas while the density of \( O_3 \) quickly saturates in He and Ar. It is known that humidity in a plasma reactor decreases the density of \( O_3 \). The presence of water vapor in a FW-DBD is inevitable due to the evaporation (physical and thermal) from water. For instance, the \( O_3 \) density dropped by a factor of 4 in gas phase by flowing the water film through the reactor [18]. The evolution of short and long lived species during the last three pulses and afterglow in the argon plasma are shown in figure 5(f). The long lived species such as \( H_2O_2 \) and \( O_2 \) accumulate over pulsing. The densities of \( H, O \) and \( OH \) rise and fall on a pulse-to-pulse basis due to formation and consumption through each pulse. For example, the rapid decline in the density of \( O \) atoms is dominantly due to ozone formation. (The transient in \( H \) density occurs over a period of 1.5–2 \( \mu \)s, and is not resolved in the figure.) The density of NO as an intermediate and less reactive species fluctuates nominally.

Densities of RNS in the gas phase after one cycle of plasma treatment are shown in figure 6 for argon, helium, air, oxygen and nitrogen plasmas including 2.5 s discharge and 10 s afterglow. Electron impact dissociation of \( N_2 \) (threshold energy of 12.25 eV) forms atomic nitrogen which initiates a sequence of reactions with \( N \) as pathways to RNS formation. Further RNS including \( N_2O_3 \) and \( HNO_3 \) (acids) are created at a slower rate during the afterglow. The dominant RNS is \( HNO_3 \) in helium and argon while the dominant RNS is \( N_2O_3 \) in air, oxygen, and
Figure 6. Density of gas phase RNS (reactive nitrogen species) during the first cycle (2.5 s of plasma exposure and 10 s of afterglow) for different feed gases. (a) Ar, (b) He, (c) air, (d) O₂ and (e) N₂. (f) RNS for an air feedstock are shown on an expanded time scale during the plasma period.

nitrogen plasmas. The higher rate of OH production in Ar and He plasmas contribute to acid formation whereas the higher rate of O production in air, oxygen and, to some degree, N₂ plasmas favor nitrogen oxides. The highest production of N atoms is in argon. Apart from electron impact dissociation, the chemical quenching of argon by N₂, \([\text{Ar}^* + \text{N}_2 \rightarrow \text{Ar} + \text{N} + \text{N}]\) yields more atomic nitrogen.

The trend in NO formation is different for each gas as NO has several production and consumption pathways by being the precursor of many reactions. For instance, in argon...
and nitrogen plasmas NO saturates quickly without significant accumulation while it builds up gradually in helium (more sources of NO). In air and oxygen, the production and consumption of NO occurs in a more pronounced manner from pulse-to-pulse. The density of NO is higher in argon and helium plasmas, around $1 \times 10^{13} \text{ cm}^{-3}$ because the higher production of OH contributes to more NO formation through $\text{OH} + \text{N} \rightarrow \text{H} + \text{NO}$ ($k = 4.7 \times 10^{-11} \text{ cm}^{-3}$).

On the other hand, NO has the lowest density in oxygen, $1 \times 10^{11} \text{ cm}^{-3}$, which is due to the low production of N which only slowly produces NO by reactions such as $\text{O}_2 + \text{N} \rightarrow \text{NO} + \text{O}$ ($k = 3.3 \times 10^{-12}, E_a = 31500 \text{ K}$). Another rapid source of NO production is through excited state $\text{N}_2^+$, $\text{O} + \text{N}_2^+ \rightarrow \text{NO} + \text{N}^+$ ($k = 7 \times 10^{-12} \text{ cm}^{-3}$) that is more dominant in air and oxygen plasma due to higher density of O.

In air, NO rapidly reacts to produce NO$_2$. Air plasma is the most efficient gas in the production of NO$_2$ as well as N$_2$O. N$_2$O is less reactive in the gas phase and has a small Henry’s law constant (solvation is a loss process to the gas phase species) and so it builds up over pulsing in the gas phase during 2.5 s of discharge. On the other hand, NO$_2$ generally saturates in the gas phase (with the exception of air) during pulsing and then increases in density in the afterglow (again air is the exception). NO$_2$ in the gas phase reaches a steady state being the most stable RNS. By the end of the afterglow (12.5 s), the density of NO$_2$ is around $1 \times 10^{12} \text{ cm}^{-3}$ in argon, helium, and air plasmas and an order of magnitude lower in oxygen and nitrogen plasmas.

NO$_2^-$ and NO$_3^-$ are short lived ions in the gas phase which are produced and consumed from pulse-to-pulse. Both ions are formed over the course of multiple pulses through several pathways. The density of NO$_3^-$ is higher than NO$_2^-$ in nitrogen, air and oxygen plasmas while it is slightly lower in helium and argon plasma. NO$_2^-$ is depleted with high O$_3$ densities by conversion to NO$_3^-_{}$:

$$\text{O}_3 + \text{NO}_2^- \rightarrow \text{NO}_3^- + \text{O}_2. \quad (6)$$

The density of HNO$_2$ in air and oxygen plasmas is the same order as the density of NO$_2$ while its density in helium, argon and nitrogen plasmas exceeds that of NO$_2$. In addition to reactions between NO and OH, HNO$_2$ is produced by associative charge exchange between HNO$_3$ + NO$_7^-$ and 3-body ion-ion recombination reaction, $\text{OH}^- + \text{NO}^+ + \text{M} \rightarrow \text{HNO}_2 + \text{M}$. The density of HNO$_2$ is the highest in argon plasmas, $6 \times 10^{14} \text{ cm}^{-3}$, followed by helium, $3 \times 10^{14} \text{ cm}^{-3}$ and air $2 \times 10^{14} \text{ cm}^{-3}$. These trends are due to the higher densities of OH$^-$ (figures 4(a) and (b)) and OH (figures 5(a) and (b)) in these plasmas. In nitrogen and oxygen plasmas the density of HNO$_2$ is approximately $2 \times 10^{13} \text{ cm}^{-3}$. HNO$_2$ is the dominant HNO in all gases.

After 2.5 s of plasma discharge, slightly more than 1500 pulses, power is turned off and simulation of the afterglow continues for 10 s, which represents one cycle of plasma water film treatment. After the discharge is extinguished, electrons and gas phase ions recombine in tens of milliseconds. Distinct phases of decay during the afterglow can be seen in figures 5 and 6. The density of short lived species such H, O, and N decrease immediately after the discharge, transient species such as OH, NO, and HNO$_2$ decay relatively fast with a timescale of hundreds of milliseconds while more stable species last longer in the gas phase during the afterglow. For the 10 s of afterglow, species are exhausted out of the reactor with the gas flow which leads to 3 orders of magnitude density reduction with the residence time of 1.6 s.

Another mechanism for loss of gas phase species is the diffusion and solvation onto the water layer. Although this process is always occurring, it is particularly important to densities of gas phase species during the afterglow when the production rate of reactive gas phase species is small. O$_3$, H$_2$, and N$_2$O are less reactive and quickly reach an equilibrium with their liquid counterparts due to their small Henry’s law constants. Therefore, these species are pumped out of the reactor leading to around 3-log density reduction without further solvation into the water or dissociation into other species.

More reactive species have a shorter lifetime during afterglow as they undergo gas phase reactions that accelerate their loss on shorter time scales than solvation into water or flow out of the reactor. These species include H$_2$O$_2$, HO$_2$, HNO$_3$, and NO$_3$. For instance, H$_2$O$_2$ loss is enhanced by dissociation reactions such as $\text{OH} + \text{H}_2\text{O}_2 \rightarrow \text{H}_2\text{O} + \text{HO}_2$ ($k = 4.53 \times 10^{-12} \text{ cm}^{-3} \text{s}^{-1}, E_a = 289 \text{ K}$) on the timescale of seconds which contributes to HO$_2$ formation. As shown in figure 5(a), after around 6 s (the time varies for each gas), H$_2$O$_2$ reaches a steady state (i.e. production equals loss). For HNO$_2$ and HNO$_3$, loss processes have shorter times than the flow time. For these species, their large Henry’s law constants accelerate their losses by solvation. (The liquid is not saturated and so solvation continues throughout the afterglow.) The density of NO$_2$ increases at the end of the discharge because its production rate is larger than its loss through the formation reactions (7a)–(7c) and its density remains unchanged during afterglow:

$$\text{OH} + \text{NO}_3 \rightarrow \text{HO}_2 + \text{NO}_2, \quad (k = 2 \times 10^{-11} \text{cm}^3\text{s}^{-1}) \quad (7a)$$

$$\text{HO}_2 + \text{NO}_3 \rightarrow \text{OH} + \text{NO}_2 + \text{O}_2, \quad (k = 4 \times 10^{-12} \text{cm}^3\text{s}^{-1}) \quad (7b)$$

$$\text{HO}_2 + \text{NO} \rightarrow \text{OH} + \text{NO}_2, \quad (k = 3.6 \times 10^{-12} \text{cm}^3\text{s}^{-1}, E_a = 270 \text{ K}) \quad (7c).$$

3.3. ROS and RNS in the liquid phase

The density ROS and RNS in the water are shown in in figures 7 and 8 for each gas mixture. Aqueous species are denoted by the ‘aq’ subscript. In our model, positive ions from the gas phase solvate in the water and undergo charge exchange to form H$_2$O$^+\text{aq}$, in which H$_2$O$^+\text{aq}$ quickly produces hydronium (H$_3$O$^+\text{aq}$) and OH$\text{aq}$ by reactions with H$_2$O$\text{aq}$. Water dissociation can occur by excitation transfer processes where excited states in the gas phase quench on the surface of
the water, $M^* + H_2O_{aq} \rightarrow M_{aq} + H_{aq} + OH_{aq}$. Electrons diffusing into the water quickly solvate, and charge transfer with $O_{2aq}$ to form $O_{2aq}^-$. Neutral species can solvate from the gas phase into water until they reach the equilibrium density with their gas phase counterpart. In this global model, water is represented as a well-stirred liquid which does not resolve diffusion inside liquid. Consequently, it is possible to overestimate solvation for species with very small Henry’s law constants. These species may, in fact, saturate at the surface of the liquid before transport deeper into the liquid, thereby reducing the solvation of gas phase species. With significant convection in the liquid (approaching the well-stirred reactor limit), the

**Figure 7.** Density of liquid phase ROS (reactive oxygen species) during the first cycle (2.5 s of plasma exposure and 10 s of afterglow) for different feed gases. (a) Ar, (b) He, (c) air, (d) $O_2$ and (e) $N_2$. 
Figure 8. Density of liquid phase RNS (reactive nitrogen species) during the first cycle (2.5 s of plasma exposure and 10 s of afterglow) for different feed gases. (a) Ar, (b) He, (c) air, (d) O\textsubscript{2} and (e) N\textsubscript{2}.

surface resident aqueous species are mixed into the volume faster than by diffusion alone.

OH is short lived in the gas phase as well as in liquid, and so the solvation of OH from gas phase should be at the proximity of the gas-liquid interface. OH\textsubscript{aq} is produced by several mechanisms, including solvation of OH from the gas phase, dissociative excitation of H\textsubscript{2}O\textsubscript{aq} by excited states and ions incident from the gas phase forming water ions. Virtually all ionization processes of H\textsubscript{2}O\textsubscript{aq} rapidly produce OH\textsubscript{aq} through the formation of hydronium: H\textsubscript{2}O\textsuperscript{+}\textsubscript{aq} + H\textsubscript{2}O\textsubscript{aq} \rightarrow H\textsubscript{3}O\textsuperscript{+}\textsubscript{aq} + OH\textsubscript{aq}. Reaction of O\textsubscript{aq} with water produces OH\textsubscript{aq}, O\textsubscript{aq} + H\textsubscript{2}O\textsubscript{aq} \rightarrow 2OH\textsubscript{aq} (k = 2.2 \times 10^{-17} \text{ cm}^{-3} \text{s}^{-1}) that
contributes to OH\textsubscript{aq} formation during afterglow, particularly in air and oxygen plasma which have a slower OH\textsubscript{aq} decay. In addition, decomposition of peroxyacetic acid (ONOOH\textsubscript{aq}) under acidic conditions yields OH\textsubscript{aq}. The density of O\textsubscript{aq} is higher with oxygen plasmas (1 \times 10^{10} \text{ cm}^{-3}), 2 times higher than that with air plasmas, and more than 3 times that with argon plasmas. Premixed dimethyl sulfoxide DMSO in water was used in the experiments as a probe to quantitatively measure OH\textsubscript{aq} \cite{6, 18}. Reaction of OH\textsubscript{aq} with DMSO produces methane sulfonic acid (CH\textsubscript{3}SOOH\textsubscript{aq}) and methyl radicals (CH\textsubscript{3}\textsuperscript{•}) that can be measured with chromatography. This measurement is akin to a titration experiment. The molarity of DMSO is chosen to be large enough so that OH\textsubscript{aq} reacts with DMSO before reacting with other species. The final densities of the reaction products of OH\textsubscript{aq} and DMSO are therefore an integral of all OH\textsubscript{aq} produced in the system. Kováčević \textit{et al} reported that the highest production of OH\textsubscript{aq} was obtained with oxygen plasmas and its inventory increases linearly with the residence time (number of PAW re-circulations). Since OH\textsubscript{aq} is highly reactive, during pulsing in the absence of DMSO, its density is in a quasi-steady state—generated and consumed with each pulse. OH\textsubscript{aq} is consumed and its density decreases continuously during the afterglow.

H\textsubscript{2}O\textsubscript{2aq} is not as reactive as OH\textsubscript{aq} due to its lower oxidation potential and is one of the most stable ROS in water. Its contribution to water purification is then more indirectly through decomposition into OH\textsubscript{aq} in which metallic catalysts such as Fe\textsuperscript{2+}, Cu\textsuperscript{2+}, and nano-TiO\textsubscript{2} are added in a solution to catalytically increase H\textsubscript{2}O\textsubscript{2aq} decomposition \cite{24}. Two major pathways to H\textsubscript{2}O\textsubscript{2aq} are solvation of H\textsubscript{2}O\textsubscript{2} (Henry’s constant 1.92 \times 10^{6}) from the gas phase or formation in the liquid through:

\[
\text{OH}^{\text{aq}} + \text{OH}^{\text{aq}} \rightarrow \text{H}_2\text{O}^{2\text{aq}}, \quad (k = 1.7 \times 10^{-11} \text{ cm}^3\text{s}^{-1}). \tag{8}
\]

The density of H\textsubscript{2}O\textsubscript{2aq} increases during pulsing and slowly reaches its Henry’s law equilibrium with the gas phase then remains unchanged during the afterglow. The most efficient plasma for H\textsubscript{2}O\textsubscript{2aq} production is sustained in argon, with the density of 5 \times 10^{16} \text{ cm}^{-3} at 10 s of afterglow. In ascending order, helium and oxygen plasmas then produce the next highest densities of H\textsubscript{2}O\textsubscript{2aq}. These trends correlate well with the density of H\textsubscript{2}O\textsubscript{2} in the gas phase suggesting that solvation may be the dominant pathway for the production of H\textsubscript{2}O\textsubscript{2aq} in this system. With helium plasmas, there is a small decrease in the H\textsubscript{2}O\textsubscript{2aq} density during the afterglow, which is due to the higher density of H2aq (see figure 5(b)) that dissociates H\textsubscript{2}O\textsubscript{2aq} by H\textsubscript{2aq} + H\textsubscript{2}O\textsubscript{2aq} \rightarrow H^{aq} + OH^{aq} + H\textsubscript{2}O\textsubscript{aq} with H\textsubscript{2}O\textsuperscript{2aq} in the water is mostly solvated from the gas phase due to its high Henry’s constant and so its density grows slowly. The solvated density of HO\textsubscript{2aq} follows that of its gas phase partner producing the maximum aqueous density with argon plasmas (1 \times 10^{14} \text{ cm}^{-3}). The higher H\textsubscript{2}O\textsubscript{2aq} density with argon and helium plasmas (approximately 2 orders of magnitude higher than in other gases) produces higher densities of HO\textsubscript{2aq} by the reaction O\textsubscript{2aq} + H\textsuperscript{aq} \rightarrow HO\textsubscript{2aq}. Although the minimum concentration of HO\textsubscript{2aq} occurs in oxygen plasmas, the lowest gas phase density of HO\textsubscript{2} occurs with nitrogen plasmas.

Hydronium, H\textsubscript{3}O\textsuperscript{+aq}, is formed by the reactions of H\textsuperscript{+aq} and H\textsubscript{2}O\textsuperscript{+aq} with water, and by hydrolysis of HNO\textsubscript{2aq} species. The density of H\textsubscript{3}O\textsuperscript{+aq} increases over time, thereby decreasing the pH of the water. The highest density of H\textsubscript{3}O\textsuperscript{+aq} and the lowest pH occurs with argon plasmas. Based solely on initial solvation of charge species, electron solvation dominantly produces O\textsubscript{2aq} whereas positive ion solvation produces H\textsubscript{3}O\textsuperscript{+aq}. Therefore, a significant imbalance between O\textsubscript{2aq} and H\textsubscript{3}O\textsuperscript{+aq} would indicate production of H\textsubscript{3}O\textsuperscript{+aq} by other processes. The major source of H\textsubscript{3}O\textsuperscript{+aq} other than positive ion solvation is hydrolysis of HNO\textsubscript{3} gas phase species, ultimately producing H\textsubscript{3}O\textsuperscript{+aq} and NO\textsubscript{3}−aq.

In the falling water experiments, Kováčević \textit{et al} measured H\textsubscript{2}O\textsubscript{2aq} in both water and DMSO solutions, where in the latter OH\textsubscript{aq} is scavenged by DMSO \cite{18}. It was shown that there is no H\textsubscript{2}O\textsubscript{2aq} in DMSO solutions when treated by air, nitrogen, and argon plasmas which implies that the origin of H\textsubscript{2}O\textsubscript{2} in its aqueous form is through the solvation of H\textsubscript{2}O\textsubscript{aq} + OH\textsubscript{aq} reaction in these systems. In helium and oxygen plasmas, solvation from the gas phase was reported to be the main pathway by observing H\textsubscript{2}O\textsubscript{2aq} in DMSO solution \cite{18}. Other studies have also reported that the scavenging of OH\textsubscript{aq} resulted in a lower H\textsubscript{2}O\textsubscript{2aq} density, a measure of the contribution of OH\textsubscript{aq} recombination pathway in the formation of hydrogen peroxide in \cite{14}. In contrast, measurement of OH\textsubscript{aq} and H\textsubscript{2}O\textsubscript{2aq} in water treated by plasma jets suggests that H\textsubscript{2}O\textsubscript{2aq} was not formed in the liquid, but instead is formed in the gas phase and then solvates into the liquid \cite{34}. However, the SVR in this FW-DBD system is significantly larger than plasma jets and the gas diffusion length to the liquid is shorter. These conditions may facilitate the solvation of more OH into the liquid which then contributes to in-liquid formation of H\textsubscript{2}O\textsubscript{2aq}. The difference between the solvated OH\textsubscript{aq} for plasmas sustained in different gas mixture in this model is nominal compared to that of H\textsubscript{2}O\textsubscript{2aq}. For example, there is a factor of 15 difference between H\textsubscript{2}O\textsubscript{2aq} produced by argon and air plasmas at the end of the discharge.

Ozone densities in the gas phase are highly dependent on the O\textsubscript{2} density in the gas mixture and the dissipated power \cite{35}. Results for gas phase ROS in figure 5 show that the highest density of O\textsubscript{3} is in oxygen plasma followed by air. The density of solvated O\textsubscript{3aq} produced by different plasmas closely follows its density in the gas phase. Other than solvation of O atoms followed by reaction with dissolved oxygen, there are few ways to produce O\textsubscript{3aq} other than solvation of O\textsubscript{3} from the gas phase. The highest O\textsubscript{3aq} density is produced by the oxygen plasma, being 2 \times 10^{16} \text{ cm}^{-3} by the end of the discharge. From high to low O\textsubscript{3aq} densities, the rankings are air, nitrogen, helium, and argon plasmas. Kováčević \textit{et al} reported that aqueous ozone densities were higher with oxygen plasma followed by air, which agrees with the results from this model \cite{18}. However the magnitudes of O\textsubscript{3aq} between this model and the experiment are quite different. These differences result from desolvation from the thin film in the model, whereas the experiment collects the activated liquid between cycles decreasing its SVR. These discrepancies will be discussed below.
These results confirm that solvation of O$_3$ from the gas into liquid is the dominant source of O$_3^{−}$aq for these conditions. O$_3^{−}$aq under basic conditions decomposes via a series of chain reactions and produces OH$\equiv$ radicals in water in which the presence of H$_2$O$_{aq}$ accelerates its decomposition via 2O$_3^{−}$aq + H$_2$O$_{aq}$ + OH$^−$ → O$_3^{−}$aq + HO$_2$aq + 2O$_2$aq + OH$^−$aq. These densities decrease during the afterflow with argon and argon plasmas compared to oxygen and air plasmas. As shown in figure 7, these results confirm that solvation of O$_3$ is larger than that of convective flow out of the reactor. HNO$_3$ has a Henry’s law constant, its rate of decay decreases by 3 orders of magnitude in all gases due to increased surface to volume ratio of the water is large and if the solvated ions are oversaturated, the species can desolvate from the liquid into the gas. This effect is most severe for species having smaller Henry’s law constants. The key to long term retention of O$_3$ in such liquids is mixing into the volume of systems with small SVRs. It was shown that the antibacterial effect of PAW strongly depends on gas-liquid mixing following plasma treatment [25]. The dynamics of solvated ions are different with helium and argon plasmas compared to oxygen and air plasmas. As shown in figure 7, the densities of O$_3^{−}$aq and OH$^−$aq, as well as HO$_2$aq, increase during the afterglow with air and oxygen plasmas. These densities decrease during the afterflow with argon and helium plasmas. With oxygen and air plasmas, O$_2^{−}$aq quickly charge exchanges with the higher density of O$_3$aq to produce O$_3^{−}$aq (k = 2.66 × 10$^{−12}$ cm$^{−3}$ s$^{−1}$). Being highly reactive, the densities of O$_3^{−}$aq rapidly decrease during the discharge and the afterglow in all gases. Reaction of O$_3^{−}$aq with OH$_{aq}$ results in the further increase in O$_2^{−}$aq and HO$_2$aq densities through reaction (9a) (k = 1.4 × 10$^{−11}$ cm$^{−3}$ s$^{−1}$). In addition, reaction of O$_3^{−}$aq with H$_2$O$^{+}$aq (equation (9b)) contributes to OH$\equiv$ formation which is more pronounced in oxygen and air plasmas by having more prolonged production of OH$_{aq}$.

\[
O_3^{−}aq + OH_{aq} → O_2^{−}aq + HO_2aq,
\]

(9a)

\[
O_3^{−}aq + H_2O^{+}aq → O_2^{−}aq + H_2O_2aq + OH_{aq}.
\]

(9b) HNO$_3$ solvates into water from the gas phase during the afterglow. Due to its large Henry’s constant, its rate of decay is larger than that of convective flow out of the reactor. HNO$_3$, with a Henry’s law constant in excess of 10$^{16}$ does not equilibrate during the afterglow with a monotonically decreasing density. HNO$_3$, with $h$ ≈ 10$^{8}$ does begin to equilibrate with HNO$_2$aq during the discharge and afterglow. The densities of HNO$_2$aq (pK$_a$ = 3.4) decrease due to hydrolysis reactions (equation (10a)) during the afterglow to reach a steady state in argon, helium, and air plasmas. Stability of weak acids such as HNO$_2$aq is pH dependent in which it dissociates to its anionic form (NO$_2^{−}$aq) when the pH is above its pK$_a$. So, the decay of HNO$_2$aq is faster in oxygen and nitrogen plasmas due to higher pH of PAW. Similarly, HNO$_3$aq is dissociated by the hydrolysis reaction (equation (10b)) that accumulates NO$_3^{−}$aq which is the dominant nitrogen-based ion in our system while the NO$_2^{−}$aq density is significantly lower. The highest density of gas phase HNO$_3$ occurs in argon plasmas, that results in the highest density of NO$_3^{−}$aq, while nitrate is an order of magnitude lower in oxygen and nitrogen plasmas due to the lower production of gas phase HNO$_3$. In our system, the products of HNO$_3$aq hydrolysis are the major source of acidification.

HNO$_2$aq + H$_2$O$_{aq}$ → NO$_2^{−}$aq + H$_2$O$_{aq}^{+}$, (10a)

HNO$_3$aq + H$_2$O$_{aq}$ → NO$_3^{−}$aq + H$_2$O$_{aq}^{+}$, (10b)

Other sources of HNO$_3$aq are the acid formation reactions from N$_2$O$_3$aq:

\[
NO_2aq + NO_2aq + H_2Oaq → HNO2aq + HNO3aq,
\]

(11a)

\[
NOaq + NO2aq + H_2Oaq → HNO2aq + HNO2aq + HNO2aq,
\]

(11b)

\[
NO2aq + Haq → HNO2aq,
\]

(11c)

\[
NO2aq + OHaq → HNO3aq,
\]

(11d)

\[
N_2O_3aq + H_2Oaq → HNO_3aq + HNO_2aq.
\]

(11e)

The density of NO$_2$aq, similar to that of ozone, quickly reaches the steady state with its counterpart in the gas phase. This equilibrium of NO$_2$aq helps maintain the density of HNO$_2$aq in spite of the latter’s rapid hydrolysis. Formation of ONOOH$_2$aq (pK$_a$ = 6.8) is pH dependent and is formed through different reactions including NO$_2$aq + OH$_{aq}$ and N$_2$O$_3$aq + H$_2$O$_{aq}$. The latter reaction has equal branchings to form HNO$_3$aq and ONOOH$_2$aq [26]. The longevity of NO$_2$aq helps buoy its density. Its conjugate base, ONOO$^{−}$aq, quickly reverts back to ONOOH$_2$aq during pulsing and the afterglow as the pH drops below 6.8. Production of ONOOH$_2$aq is higher with air plasmas (1 × 10$^{15}$ cm$^{−3}$) due to the higher and prolonged density of OH$_{aq}$ and a high concentration of N$_2$O$_3$. Furthermore, the decay of ONOOH$_2$aq (less in PAW produced by air plasmas than other gases) into NO$_3^{−}$aq, H$_2$O$_{aq}^{+}$, and NO$_2$aq prolongs the lifetime of these species during the afterglow. Chemical analysis of phenol in PAW by Lukes et al., and the formation of hydroxylated and nitrated products of phenol, indicated the decomposition of ONOOH$_2$aq and post-discharge production of OH$_{aq}$ and NO$_2$aq [36]. Peroxynitric acid (H$_2$O$_2$NO$_2$aq) is formed in the gas phase through 3-body reactions and solvates with a large Henry’s law constant. Its density in the gas phase as well as in liquid is higher in air and argon plasmas. Although H$_2$O$_2$NO$_2$aq decays thermally to H$_2$O$_{aq}$ + NO$_2$aq and HNO$_2$aq, it is relatively stable in helium, argon, and air produced PAW up to 10 s afterglow. The hydrolysis of H$_2$O$_2$aq (pK$_a$ = 4.8), particularly in helium and argon.
produced PAW, contributes to further increase in the density H$_2$O$_{aq}$^− and O$_2$^− $aq$. HO$_{aq}$ also reacts with OH$_{aq}$ to form O$_{aq}$ and H$_2$O$_{aq}$.  

4. Re-circulating PAW

To model the recirculation of the PAW, the densities after 12.5 s from all gas and liquid species were used as the input density for the simulation of the next cycle, and this plasma treatment was repeated for ten cycles as in the experiment. In the experiment, PAW was collected in a container at the end of one cycle and then was loaded again into the DBD reactor for the next rounds of plasma treatments. Therefore, the evolution of liquid phase species in the experiment can be affected by more complicated environmental factors such as mixing, exposure to the ambient air, and changes in the surface-to-volume ratios. In the model, the time interval between each re-circulation was estimated to be 10 s. During all of this time, the SVR was the same as during the plasma activation.

In the experiment, DMSO was added to the water prior to the plasma treatment. As a result, OH$_{aq}$ was rapidly titrated out of the water, producing a time integrated value for OH$_{aq}$ production. The rapid titration of OH$_{aq}$ decreases the rates of reactions that are initiated by OH$_{aq}$. To compare the computed densities of OH$_{aq}$ with measurements [18] the computed density of OH$_{aq}$ was integrated over the total time (12.5 s) of the process and multiplied by the lifetime of OH$_{aq}$ in our system (3 ms) to compensate the quick reaction of OH$_{aq}$ with the molecular probe (DMSO). The time integrated concentration of OH$_{aq}$ is shown in figure 9 for the first cycle. The OH$_{aq}$ production in water is approximately the same during all cycles, suggesting a linear growth over the 10 cycles of recirculation. The highest integrated density of OH$_{aq}$ is in air plasma produced PAW (20 $\mu$M) due to the decomposition of species such as peroxinitrous acid and ozone (O$_{aq}$ = H$_2$O$_{aq}$ + H$_2$O$_{aq}$ → O$_{aq}$ + OH$_{aq}$ + H$_2$O$_{aq}$). Considering a linear growth over 10 cycles and multiplying the data in figure 9, the density of OH$_{aq}$ in our model is in agreement with the concentration, ∼200 $\mu$M, after the last cycle measured in the experiment. The density of OH$_{aq}$ measured in the experiment was approximately 100 $\mu$M in nitrogen, argon, and helium. The lowest integrated OH$_{aq}$ density as shown in figure 9 is with the argon plasma as 5 $\mu$M after one cycle.

Densities of H$_2$O$_{aq}$ in PAW produced by at the end of each cycle (12.5 s) are shown in figure 10(a). The highest density of H$_2$O$_{aq}$ occurs with argon plasmas (plotted on the right axis), 0.25 mM after the 10th cycle, which correlates well with the experimental data. Kovacević et al showed that the highest production rate of H$_2$O$_{aq}$ in water is in argon plasma and the highest concentration is around 0.6 mM after the 10th cycle [18]. Both the simulation and the experiment indicate that H$_2$O$_{aq}$ accumulates over recirculation though the trend is different for each gas. With argon, helium, and nitrogen plasmas, there are incremental increases in the density of H$_2$O$_{aq}$ with each cycle, whereas with air and oxygen plasmas H$_2$O$_{aq}$ reaches a near steady state. According to the experiment, the lowest density of H$_2$O$_{aq}$ was obtained with oxygen and nitrogen plasmas while the density of H$_2$O$_{aq}$ was a factor of 3 higher with air plasma. As shown in figure 5, production of H$_2$O$_{aq}$ in the gas phase during the discharge is also lower in air and nitrogen plasmas, which correlates well with its solvated counterpart suggesting solvation is the dominant pathway.

Densities of O$_{aq}$ at the end of the afterglow for each cycle are shown in figure 10(b). The concentration of O$_{aq}$ is approximately 21 nM and 12 nM when using oxygen and air plasmas, respectively, higher than in other gases. The density of O$_{aq}$ is lower by 2 orders of magnitude when using helium and argon plasma. There is no significant increment in O$_{aq}$ density with re-circulation. These trends are attributed to the gas phase density of O$_3$ in this system reaching a steady state over the course of plasma pulses and O$_{aq}$ saturating by having small Henry’s constant. Therefore, the concentration of O$_{aq}$ will not further increase with increasing the residence time or cycles. The lower concentrations of O$_{aq}$ when using He, Ar and N$_2$ plasmas is a consequence of the only source of oxygen being either impurities or dissociation of water vapor. The predicted ranking of densities of O$_{aq}$ for each gas correlates well with the results of the experiment—the highest O$_{aq}$ occurs with oxygen plasmas, followed by air and nitrogen plasmas.

Having reproduced trends of the experiments, the predicted concentrations of O$_{aq}$ presented in figure 10(b) are significantly lower than the experimental measurements. The differences can be attributed to the rapid loss of O$_{aq}$ in our model due to desolvation during the afterglow, as shown in figure 7. In the model, the thin-film nature of the plasma-water system is retained during the afterglow between plasma treatment cycles. With gas phase O$_3$ convected away from...
the liquid during the afterglow, the Henry’s law equilibrium concentration of $O_{3aq}$ also decreases. In the experiment, mixing, collection of the water into a volume having a smaller surface-to-volume ratio, and environmental factors can reduce the desolvation rate of $O_{3aq}$ resulting in retaining a higher concentration of dissolved ozone between cycles.

With these considerations, the predicted densities of $O_{3aq}$ at the end of the plasma discharge (and before the rapid desolvation during the afterglow) are perhaps the better comparison to the experiment. The predicted $O_{3aq}$ at the end of plasma pulsing in oxygen (the highest density) is approximately 20 µM while in the experiment the concentration was 8 µM. For plasmas sustained in nitrogen, the experimental density of $O_{3aq}$ is $\sim$2 µM which is underestimated by the simulation as 0.15 µM. The concentrations of $O_{3aq}$ for air plasmas is 5.2 µM in the experiment and 12 µM in the model. In the the experiment, the density of $O_{3aq}$ for plasmas in helium fluctuates around 0.1 µM without a significant increase in density with recirculation, similar to the simulation results which also records a concentration of 0.1 µM. These comparisons between the experimental measurements of $O_{3aq}$ and the simulation results at the end of the pulsing qualitatively agree, and emphasize the need to manage the desolvation of species from thin water films.

The accumulation of nitrate, $NO_3^{-aq}$, in PAW for plasmas sustained in different gases at the end of each cycle (discharge and afterglow) is shown in figure 10(c). In all gases, the $NO_3^{-aq}$ density increases linearly with the cycles of recirculation (residence time). The highest predicted concentrations of nitrate are 1.2 mM in argon and 0.45 mM in air plasmas. The lowest densities of nitrate are in oxygen and nitrogen plasmas 0.04 mM. In the experiment, the densities of $NO_3^{-aq}$ in descending order were in air (2.5 mM), nitrogen (0.64 mM), and argon (0.22 mM) plasmas, while the density was nominal in oxygen and helium plasmas. These comparisons are complicated by variations in power deposition in the experiment, while power was constant across gases in the model. When running isolated cases with powers aligned with the experiments, the predicted density of $NO_3^{-aq}$ is highest in air plasmas (instead of argon), as in the experiment.

Solvation of $NO_x$ species from the gas and the dissociation of acids in the water generate $H_3O^{+aq}$ and decrease the pH of the PAW. Since $HNO_3aq$ is a strong acid its hydrolysis (forming...
NO$_3^{-aq}$ is the major source of H$_3$O$^+aq$. Therefore, the trend of pH in PAW follows the NO$_3^{-aq}$ density as shown in figure 10(d). The pH in a descending order at the last cycle is 4.4 in oxygen and nitrogen, 3.6 in helium, 3.3 in air, and 2.9 in argon plasmas. The lowest pH in experiments was in air plasmas followed by nitrogen. When running powers identical to the experiments, the predicted pH is also lowest for air plasmas.

5. Concluding remarks

Production of PAW not only depends on the formation of gas phase precursors but also their transport and subsequent solvation into the water. A computational investigation was performed of the formation of PAW in a FW-DBD in which a thin film of water is treated through serveral cycles while varying the feedstock gas (Ar, He, N$_2$, O$_2$, air) [18]. In each case, the feedstock gases become saturated with water vapor, and so precursor formation was in part determined by the reactions of the plasma activated feedstock gases with the water vapor. For example, the most efficient gases for OH production in the gas phase were Ar and He, in part due to these interactions.

Although OH$_{aq}$ is probably the most important oxidant in water treatment, it has a short lifetime in water, being produced and consumed during each discharge pulse and each cycle. The major product of OH$_{aq}$ reactions, H$_2$O$_{2aq}$, is more stable and accumulates during the early cycles of cyclic treatment of PAW in the FW-DBD, with the highest production rate being in argon plasmas. Prolonged densities of OH$_{aq}$ result from in liquid reactions of O$_3^{-aq}$ and ONOOH$_{aq}$. Nitrate (NO$_3^{-aq}$), produced dominantly by hydrolysis of HNO$_3$ gas phase species, accumulates with successive cycles, the largest densities being with Ar and air discharges, leading to these systems also being the most acidic due to the co-production of H$_3$O$^+aq$ during the hydrolysis.

In general, the trends in the computed results for components in the PAW as a function of gas mixture aligned with experiments by Kovačević et al [18], which lends credence to the calculations. The differences can in large part be attributed to the use of a global model for a geometrically complex system and uncertainties in power deposition. The exception to this agreement is in the measured and computed values of O$_2^{aq}$, and the source of that disagreement is a major lesson of this work. In both experiment and computations, the concentrations of O$_2^{aq}$ saturated, a consequence of the low Henry’s law constant of O$_2$. However, the predicted densities of O$_2^{aq}$ at the end of after glow cycles were significantly below those of the experiment. The explanation is likely due to the manner in which the PAW was treated between plasma cycles. In the computations, the system was maintained in a thin-film configuration with a large SVR, which enabled the O$_2^{aq}$ to desolvate. In the experiment, the collection and recirculation of the PAW produce mixing and resulted in a smaller SVR which reduced the rate of de-solvation of O$_3^{aq}$. When comparing the post-plasma, saturated concentrations of O$_3^{aq}$ prior to desolvation, the agreement between calculations and experiments was reasonable. These results emphasize the need to actively manage the PAW to prevent (or enhance) the rate of desolvation of species [37].

Modeling the many thousands of pulses and tens of seconds of processing time for five gas mixtures was a computationally intensive task, which would be beyond the current capabilities of the 2-dimensional or 3-dimensional models required to address, in detail, a geometrically complex system. Questions posed by a referee in this regard include whether such modeling can be predictive? The direct answer is that global modeling can be predictive, as demonstrated here, in a semi-quantitative way. However, as the geometric complexity of a system increases, the expectations should decrease that global modeling can provide a side-by-side quantitative representation of the experiment. The value of global modeling in this regard is in emphasizing important mechanisms and trends.

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