J. Phys. D: Appl. Phys. 47 (2014) 475203 (11pp)

# Atmospheric pressure plasma jets interacting with liquid covered tissue: touching and not-touching the liquid

#### Seth A Norberg<sup>1</sup>, Wei Tian<sup>2</sup>, Eric Johnsen<sup>1</sup> and Mark J Kushner<sup>3</sup>

<sup>1</sup> Department of Mechanical Engineering, University of Michigan, 2350 Hayward Street, Ann Arbor, MI 48109-2125 USA

<sup>2</sup> Department of Nuclear Engineering and Radiological Science, University of Michigan, 2355 Bonisteel Boulevard, Ann Arbor, MI 48109-2104 USA

<sup>3</sup> Department of Electrical Engineering and Computer Science, University of Michigan, 1301 Beal Avenue, Ann Arbor, MI 48109-2122, USA

E-mail: norbergs@umich.edu, ejohnsen@umich.edu, bucktian@umich.edu and mjkush@umich.edu

Received 21 July 2014, revised 7 September 2014 Accepted for publication 26 September 2014 Published 5 November 2014

#### Abstract

In the use of atmospheric pressure plasma jets in biological applications, the plasma-produced charged and neutral species in the plume of the jet often interact with a thin layer of liquid covering the tissue being treated. The plasma-produced reactivity must then penetrate through the liquid layer to reach the tissue. In this computational investigation, a plasma jet created by a single discharge pulse at three different voltages was directed onto a 200 $\mu$ m water layer covering tissue followed by a 10 s afterglow. The magnitude of the voltage and its pulse length determined if the ionization wave producing the plasma plume reached the surface of the liquid. When the ionization wave touches the surface, significantly more charged species were created in the water layer with H<sub>3</sub>O<sup>+</sup><sub>aq</sub>, O<sub>3</sub><sup>-</sup><sub>aq</sub>, and O<sub>2</sub><sup>-</sup><sub>aq</sub> being the dominant terminal species. More aqueous OH<sub>aq</sub>, H<sub>2</sub>O<sub>2aq</sub>, and O<sub>3aq</sub> were also formed when the plasma plume touches the surface. The single pulse examined here corresponds to a low repetition rate plasma jet where reactive species would be blown out of the volume between pulses and there is not recirculation of flow or turbulence. For these conditions, N<sub>x</sub>O<sub>y</sub> species do not accumulate in the volume. As a result, aqueous nitrites, nitrates, and peroxynitrite, and the HNO<sub>3aq</sub> and HOONO<sub>aq</sub>, which trace their origin to solvated N<sub>x</sub>O<sub>y</sub>, have low densities.

Keywords: plasma jet, plasmas on liquids, atmospheric pressure plasma, plasma activated water

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

#### 1. Introduction

The use of atmospheric pressure plasma jets in the context of plasma medicine for wound healing, sterilization of surfaces and cancer treatment is an increasingly important area of research [1-3]. Often the surface being treated is cellular tissue, which is covered by a thin liquid layer of a blood serum-like liquid a few hundred microns thick [4]. Instead of directly interacting with the tissue, the plasma-produced neutral reactive species, ions and photons first react with the liquid layer.

Gas phase species solvate into the liquid and undergo additional reactions before the plasma produced activation energy reaches the underlying tissue. Photolysis and photoionization from plasma-produced ultraviolet and vacuum-ultraviolet (UV/VUV) photons may directly produce active species (neutral reactive species and ions) in the top layer of the liquid that subsequently react to form other aqueous ions and neutral reactive species.

Many recent investigations have addressed the manner in which plasma jets produce reactivity in liquids, starting with water [5–7]. The majority of these studies have used a rare-gas seeded with a small amount (usually <1%) of a reactive gas (e.g.,  $O_2$ ,  $N_2$ ,  $H_2O$ ) as the media flowing through the plasma tube, which is then exhausted into room air. RONS (reactive oxygen and nitrogen species) are directly produced in the plasma jet and by mixing with the ambient air [8]. Configurations using gas shrouds around the jet have been investigated to control the mixing of the ambient air with the jet to then control reactive species production [9, 10].

A parameter in these investigations is whether the plasma plume emanating from the jet contacts the liquid layer (touching) or is not in direct contact with the liquid layer (nottouching). Experimentally, this distinction is usually made on whether the luminous visible plume from the jet extends to the surface of the liquid. Significantly different results for the reactive species composition in the liquid have been reported based on whether the plasma jet is touching or not-touching the liquid [11]. The delivery of electric fields to the underlying tissue is also expected to be sensitive to the touching or not-touching conditions. In this paper, we discuss results from a computational investigation of plasma jets incident onto a thin water layer over a tissue-like dielectric. Different applied voltages are used which result in touching and not-touching configurations. The proportion and magnitude of aqueous charged and neutral species that diffuse to the underlying tissue depends on the touching or not-touching conditions. Aqueous ions diffusing through the liquid to the underlying tissue are significantly greater for touching conditions due to the rapid solvation of gas phase ions afforded by the contact. Neutral aqueous species produced by the jet are less sensitive to the touch or no-touch conditions, and scale more proportionately with the energy deposition per pulse in the jet.

The model used in this investigation is described in section 2. A comparison of touching and not-touching plasma jets onto thin liquid layers is discussed in section 3. Our concluding remarks are in section 4.

#### 2. Description of model

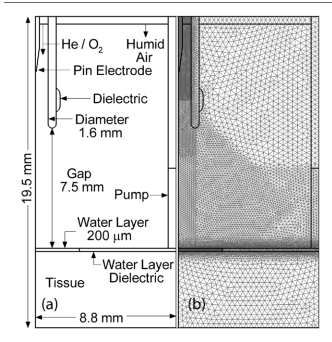
The 2D cylindrically symmetric, plasma hydrodynamics model utilized in this investigation is *nonPDPSIM* which is executed on an unstructured mesh as described in detail in [12, 13]. For the purpose of this study, nonPDPSIM consists of three major modules-the plasma transport module, the radiation transport module and the fluid transport module. In the plasma transport module, Poisson's equation and continuity equations for all charged species (in the volume and on surfaces) are integrated in time using a fully implicit Newtoniteration technique [13]. Spatial derivatives and divergence operators are expressed in fully conservative finite-volume form. Each time step is then followed by an implicit update of the electron energy equation for electron temperature, neutral species continuity equations and radiation transport. Transport coefficients and rate coefficients for electrons as a function of electron temperature are obtained from solutions of the stationary Boltzmann's for the electron energy distribution. These tabulated coefficients which are interpolated during execution of the model are updated on sub-nanosecond time scales to reflect changing species and mole fractions. Radiation transport addresses photoionization of  $O_2$  and  $H_2O$  in the gas phase, and photoionization and photo-dissociation of  $H_2O_{aq}$  in the liquid phase. (The subscript <sub>aq</sub> denotes aqueous or in liquid species.) This portion of the model is discussed in detail in [13, 14].

In the fluid module, a modified form of the Navier–Stokes equations (continuity, momentum, energy) are solved to produce the advective flow field while including body forces (e.g., momentum transfer from charged particles) and heat sources (e.g., Joule heating and change in enthalpy due to reactions) produced in the plasma transport module [15]. Since the gas constituents have large differences in atomic or molecular mass, the mass continuity equation is replaced by a number-density continuity equation and a fourth equation for the average molecular weight. These equations are simultaneously and implicitly solved using numerically derived Jacobian elements on the same unstructured mesh as the plasma calculation. The individual continuity equations for all species are simultaneously integrated using the advective flow field as a check for the average molecular weight.

The initial liquid layer in this investigation is pure water with 8 ppm dissolved  $O_2$ . The water layer is included in the model using the same algorithms as in the gas phase. In principle, the water layer is just another region of the plasma albeit with a higher density and different reaction mechanism. Surface tension is not explicitly calculated in the model we simply specify the boundary between the liquid and gas phase. However, evaporation of the water layer into the gas phase is included. The methods used to couple the gas phase plasma and the liquid phase calculations, the reaction mechanism, and method for representing water vapor evaporating from the reactive and dielectric water layers (see below) are discussed in [14].

The fluid module is first executed without the plasma to establish the advective flow field. For the geometry and flow rates of interest, the flow field is established in tens of ms of integration time using timesteps that are typically many times the Courant limit for the smallest cells in the mesh. The plasma is then initiated as described below, and the fluid and plasma modules are simultaneously integrated until the voltage pulse is terminated and space charge is allowed to dissipate. At that point, we stop solving Poisson's equation and assume local charge neutrality. The fluid module and continuity equations of individual species continue to be integrated with ns timesteps to account for rapid reactions that initially take place. As the reactivity diminishes, the timesteps grow to  $1\,\mu s$  over 50 ms of flow during which the majority of reactive gaseous species are blown out of the computational domain. At this point, the fluid module is terminated and the reactions of the aqueous species are continued to be solved with a time step of  $10-100\,\mu s$  until the computation is terminated at 10 s.

Based primarily on the dimensions of the KinPen developed by the INP Greifswald [9, 10]. The geometry used in this investigation for the plasma jet is a cylindrical tube with a diameter of 1.6 mm and walls that are 0.4 mm thick. The tip of the coaxial electrode, 0.25 mm in diameter, is 3.5 mm from

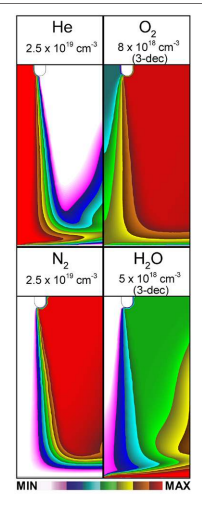


**Figure 1.** Schematic of the cylindrically symmetric model geometry. (*a*) The total computational domain is 1.95 cm in height and 0.88 cm in radius. (*b*) Numerical mesh showing different refinement zones.

the end of the tube, as shown in figure 1. In this configuration, there is an axial component of the electric field that better enables the plasma to intersect with the liquid [16, 17]. This configuration differs from those that have an electric field that is dominantly perpendicular to the gas flow and which produces a plume that is dominated by neutral reactive species [18]. The end of the tube is  $7.5 \,\mathrm{mm}$  above a  $200 \,\mu\mathrm{m}$  thick water layer covering the tissue-like dielectric. The portion of the water treated as plasma extends to a radius of 2.75 mm. For larger radii, the water is not treated as a reactive zone but rather is represented simply as a dielectric material. These separate water zones were implemented to lower the computational cost. Water evaporates from both water zones. The numerical mesh, also shown in figure 1, consists of 12700 nodes with several refinement zones having spatial resolution varying from  $24\,\mu\text{m}$  near the tip of the electrode and within the water layer to  $300 \,\mu\text{m}$  in the periphery of the domain.

Helium seeded with O<sub>2</sub> (He/O<sub>2</sub> = 99.8/0.2, 5 slm) is flowed through the tube. Humid air (N<sub>2</sub>/O<sub>2</sub>/H<sub>2</sub>O =79.5/20/0.5) flows at a lower rate outside the tube (4 slm) to entrain the helium and guide the helium flow over the surface. The resulting mixture flow out of the computational domain through a pump port on the side boundary. The material for the tube has a dielectric constant  $\varepsilon_r = \varepsilon/\varepsilon_o$ =4, the tissue underlying the liquid has  $\varepsilon_r = 5$  and the non-reactive water layer has  $\varepsilon_r = 80$ . At the beginning of the plasma period, the voltage pulse is applied for 50 ns (5 ns rise time).

A single discharge pulse of a plasma jet, its afterglow and activation of a thin water layer are discussed here. Actual systems are operated with a pulse-repetition frequency (PRF). Plasma jet treatment of tissue differs significantly from dielectric barrier discharge (DBD) treatment in many ways—one



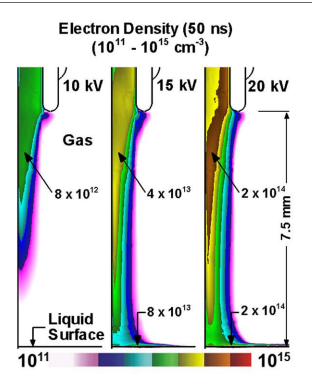
**Figure 2.** Flow profiles at 8 ms prior to initiating the discharge for (*a*) He, (*b*)  $O_2$ , (*c*)  $N_2$  and (*d*)  $H_2O$ . The water vapor is blown from the surface of the water layer by the jet. The maximum value is noted in each frame. Contours are on a linear scale unless the number of decades for a log scale is noted.

being that the gas in a plasma jet has a finite residence time in the vicinity of the tissue. Depending on flow rates, jet diameter and situational factors (e.g., turbulence, eddies produced by local structures [19]), the gas and plasma may smoothly blow out of the vicinity of the tissue in a laminar manner with air diffusing into the jet or there may be significant recirculation. All of these effects are important to the final fluence of reactants to the tissue underlying the liquid. The results of this study are therefore applicable to conditions of laminar flow with a PRF of less than a few kHz where the plasma activated gas flows out of the volume between pulses.

## 3. Touching and not-touching plasma jets onto a liquid layer

#### 3.1. Transport of gas phase RONS

The pre-pulse steady-state densities of He,  $O_2$ ,  $N_2$  and  $H_2O$  in the fluid flow-field are shown in figure 2 from the exit nozzle of the jet. Helium flows down from the tube into the ambient air where the helium is largely confined to the center of the



**Figure 3.** Electron density for applied potentials of -10kV, -15kV and -20kV at 50 ns prior to pulse termination. The electron plume for -10kV does not touch the liquid surface. The electron plume for -15kV touches at 35 ns (in contact with water and spreading for 15 ns) and the plume for -20kV touches at 20 ns (in contact for 30 ns). Contours are on a log-scale over 4 decades.

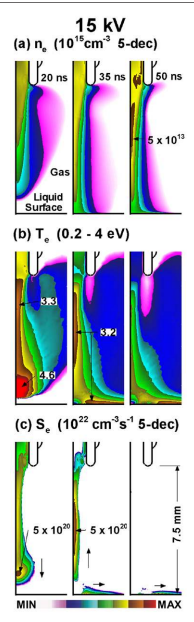
jet by air diffusing into the core. The He then flows radially outward from the stagnation point on axis. The axial speed of the flow field inside the tube is  $42 \,\mathrm{m \, s^{-1}}$ . The corresponding Reynolds number is Re = 630, which is less than the value for onset of turbulence for pipe flow, Re = 2300. So the flow should be laminar leaving the tube. Once the jet leaves the tube, for Re = 630 the onset of turbulence will occur at about  $50r_0$  from the outlet, where  $r_0$  is the jet radius [20]. For our conditions, the onset of turbulence would be expected at about 4 cm from the tube outlet, which is larger than our outletsurface distance. The increase in the He concentration at the right of the panel is a vortex at larger radii. The density of O<sub>2</sub> is shown over three decades to highlight the small amount,  $5.9 \times 10^{16}$  cm<sup>-3</sup>, contained in the input He gas flow. The diffusion of air into the He jet increases the O<sub>2</sub> concentration to  $3.9 \times 10^{17} \text{ cm}^{-3}$ , on axis 1 mm above the water. N<sub>2</sub> from the ambient is absent from the interior of the tube and becomes entrained in the He jet further from the exit as air diffuses into the jet stream. The density of N<sub>2</sub> is  $1.3 \times 10^{18} \text{ cm}^{-3}$  on axis 1 mm above the water. The water vapor that evaporates from the water layer is displaced by the helium jet and flows to the pump. The density of H<sub>2</sub>O in the ambient air is  $1.2 \times 10^{17}$  cm<sup>-3</sup>. Directly above the liquid water on the axis, the density of H<sub>2</sub>O is low,  $\approx 10^{16}$  cm<sup>-3</sup>, due to the inability for the ambient water vapor to diffuse into the core of the helium jet. The flow of the jet displaces the evaporating water vapor directly above the liquid layer at small radii and compresses the water vapor

at larger radii. The reactive species delivered to the surface of the liquid are determined by the species initially produced by electron impact excitation, dissociation and ionization in the plasma plume, their subsequent reactions, convection in the fluid flow field and density gradient driven diffusion.

Plasma jets for three voltages were investigated— - 10kV, -15kV and -20kV. The voltage pulses have a 5ns rise time and 45ns duration. A cloud of plasma with density of 10<sup>11</sup> cm<sup>-3</sup> and radius 0.7 mm is seeded at the tip of the powered electrode for ignition. The electron densities,  $n_e$ , at the end of the discharge pulse for each voltage are shown in figure 3. The electron avalanche begins as an ionization wave at the tip of the pin electrode, and quickly assumes an annular, wall hugging mode [21, 22], which charges the inside surface of the glass tube. As the ionization wave exits the tube, the avalanche returns to the axis. This is the same behavior for all voltages— $n_e$  is annular and surface-hugging inside the tube and transitions to on-axis within a few mm of exiting the tube. The  $n_e$  on axis 1 mm from the end of the tube is  $6.7 \times 10^{12}$  cm<sup>-3</sup>,  $2.7 \times 10^{13}$  cm<sup>-3</sup>, and  $6.9 \times 10^{13}$  cm<sup>-3</sup> for -10kV, -15kV and -20kV. In the -10kV case, the plasma plume, represented by the electron density, does not reach the surface of the water with a significant density and this is considered the indirect, or not-touching, case. The plasma plumes for the -15 kV and -20 kV cases both reach the water and are the direct, or touching cases. The electron density 1 mm above the water layer on axis is  $4.2 \times 10^{12}$  cm<sup>-3</sup> and  $6.7 \times 10^{12}$  cm<sup>-3</sup> for the -15 kV and -20 kV cases.

The electron density, electron temperature  $(T_e)$  and electron impact ionization source  $(S_e)$  are shown in figure 4 for the -15 kV case. The speed of the ionization wave depends on the applied voltage. For the  $-15 \,\text{kV}$  case, the electron plume touches the surface of the water at 26 ns. A conducting channel of increasing electron density is produced on axis as the capacitance of the water layer is charged until the voltage is terminated.  $T_e$ , and  $S_e$  trace the trajectory of the head of the ionization wave (the *plasma bullet*) or guided streamer [8].  $T_e$ and  $S_e$  in the head of the plasma bullet midway through the gap are 5 eV and 1  $\times$  10<sup>21</sup> cm<sup>-3</sup> s<sup>-1</sup>. The mushroom shaped luminous plasma bullet from the simulation before striking the surface of the water layer is similar to those shapes experimentally observed for plasma bullets [23]. Upon striking the water layer, a restrike, positive ionization wave propagates back up the plasma column.

The ionization wave generated in the plasma tube propagates through the helium dominated channel and, if sufficient duration or length, intersects with the liquid layer. Since the liquid layer has largely dielectric properties, the discharge at this point behaves similarly to a DBD—the plasma spreads on the surface of the liquid as the surface charges. This spreading results from many causes. The large dielectric constant of the liquid excludes electric fields, on a relative basis, out of the liquid in favor of the air. The lower electric field and higher neutral density in the water produces a low E/N [electric field/(gas or liquid number density)] that will not self-sustain the plasma. For example, at the instant the ionization wave reaches the surface of the water at 26 ns, the E/N = 60 Td (1 Td =  $10^{-17}$ V cm<sup>2</sup>) 25  $\mu$ m above the surface of the water



**Figure 4.** Plasma dynamics -15 kV. (*a*) Electron density, (*b*) electron temperature and (*c*) electron impact ionization source at 20 ns, 35 ns and 50 ns after start of the pulse. At 20 ns the ionization wave has not touched the surface. The spread of the ionization as a surface ionization wave continues after touching. Contours are on a linear scale unless the number of decades for a log scale is noted.

while the E/N = 0.0025 Td  $25 \mu$ m below the surface of the water. These latter values of E/N are well below that required to self-sustain a plasma. The charged particle inventory in the water is therefore a result of electrons and ions solvating into the liquid, photoionization and water chemistry, and not direct electron impact dissociation and ionization.

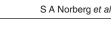
There is also *de facto* charging of the surface layer of the liquid due to the long dielectric relaxation time of the low-conductivity water. The solvation of electrons into the water, and charge exchange from positive ions with water molecules, occurs on a ps to ns timescale, transferring charge to the water. The low mobility of these charged species in the water

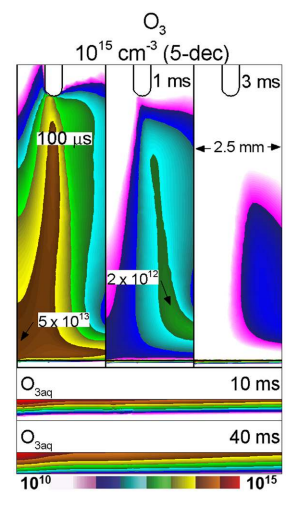
confines the transferred charge to the surface of the water for the duration of the discharge. The charge injection into the water is augmented by photoionization of the water by UV/ VUV photons produced by the plasma adjacent to the water. For these conditions, the charge at the surface of the water at the end of the discharge pulse is due 90% from solvation of gas phase charged particles into the liquid and 10% from photoionization.

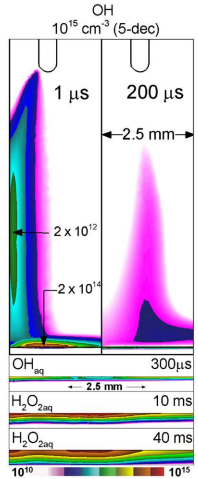
Plasma plumes that do not touch the water layer lack the nearly immediate electron solvation, charge exchange and photoionization (and photolysis) reactions at the surface of the liquid that result from plasma contact. The spreading of the plasma on the surface of the water is important to the production of charged aqueous species. The solvation of charged species is distributed over a larger radius, and the close contact enables UV/VUV radiation, that has a short mean-free-path, about  $12\mu m$ , to penetrate into the water. The electron density in the not-touching plasma plume largely decays due to recombination with molecular ions, dominantly N<sub>4</sub><sup>+</sup> and O<sub>2</sub><sup>+</sup>, and electron attachment to O2 in many to tens of microseconds. This time is short compared to the time for the convective gas flow to reach the liquid, about 0.34 ms. So the vast majority of electrons will have recombined or attached prior to reaching the liquid. The end result is that the effluent of the plume consists of a low density ion-ion plasma. Since the time scale for ion-ion recombination can be commensurate with the time for the convective gas flow to reach the liquid, a low density of ions does reach the water even in the nottouching case. For example, for the -10kV not-touching case, the maximum gas phase negative ion density adjacent to the water is  $10^{10}$  cm<sup>-3</sup> at tens of  $\mu$ s after the discharge pulse. Note that this long time scale ion-ion plume occurs for the higher voltages as well. As discussed below, its influence is simply smaller in the touching cases.

The gas dynamics play an important role in distributing gas phase reactive species to the liquid. For example, the density of  $O_3$  is shown in figure 5 for the -15 kV case. The O atoms that produce  $O_3$  by a 3-body reaction with  $O_2$  are formed dominantly inside the tube by electron impact dissociation of the O<sub>2</sub> additive. Since the density of O<sub>2</sub> is low in the tube, the formation of  $O_3$  is low. The formation of  $O_3$  is dominantly outside the tube where the  $O_2$  in the air diffuses into the plasma plume and reacts with the O atoms formed in the tube. Once formed, O<sub>3</sub> is relatively unreactive in the absence of organic impurities in the air, and so flows with the plume to the surface of the liquid. As the plume spreads along the surface of the water, the O<sub>3</sub> solvates to form the terminal species  $O_{3aq}$ . The solvation of  $O_3$  is therefore fairly uniform over a large radius. The majority of O<sub>3</sub> flows out of the computational domain. Here is where the details of the local flow dynamics are important. Vortices, turbulence or stagnationpoint flow that are produced by, for example, non-vertical jets or the structure of sample-wells will recirculate species such as O3 back to the water layer instead of directly flowing out of the region of interest.

 $O_{3aq}$  largely results from solvation of gas phase  $O_3$  and the production of  $O_3$  depends largely on the inventory of O atoms produced by the discharge. The formation of  $O_3$  is therefore







**Figure 5.** Ozone density in the gas phase and in the liquid for -15 kV at varying times after the discharge pulse. Ozone is relatively non-reactive in both phases. Contours are on a 5-decade log-scale.

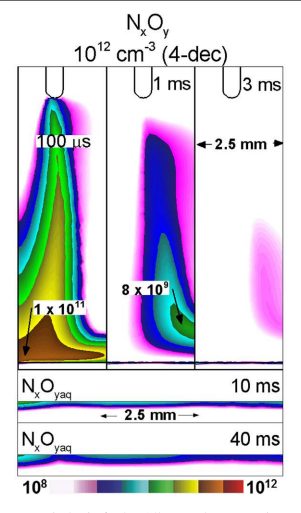
only weakly dependent on whether the plasma touches or does not tough the liquid, other than plasma touching plumes are typically of longer duration and so produce more O atoms in the vicinity of the liquid. The maximum O<sub>3</sub> density 1 mm above the liquid layer at 100 $\mu$ s is 1.1 × 10<sup>13</sup> cm<sup>-3</sup>, 4.6 × 10<sup>13</sup> cm<sup>-3</sup>, and 1.4 × 10<sup>14</sup> cm<sup>-3</sup> for -10 kV, -15 kV and -20 kV. The end result is that O<sub>3aq</sub> is produced in large amounts in both the touching and not-touching cases.

The production of the hydroxyl radical, OH, in the gas phase significantly differs from that of O<sub>3</sub>. OH is initially formed by electron impact dissociation of H<sub>2</sub>O during the interaction of the plasma and the humidity in the air and, more importantly, the evaporated water vapor above the water layer. The density of OH is shown in figure 6 at 1 $\mu$ s and 200 $\mu$ s after the discharge pulse for the -15 kV case. OH is largely produced in the plasma column by electron impact dissociation of water diffusing into the jet, generating a maximum density of 2.2 × 10<sup>12</sup> cm<sup>-3</sup> at mid-gap at the end of the pulse. A larger density is produced adjacent to the water by the spreading plasma over the liquid intersecting with the evaporating water vapor. The maximum OH density adjacent to the water is 7.2 × 10<sup>14</sup> cm<sup>-3</sup> at the end of the discharge pulse. Although the production of OH does not significantly change in mid-gap

**Figure 6.** OH in the gas phase and water, and  $H_2O_2$  in the water for the -15 kV case at varying times after the discharge pulse. At  $1\mu$ s, the highest OH concentration is adjacent to the water layer where the electron plume intersects the evaporating water. Over the next  $200\mu$ s the OH reacts or blows away. The evolution of  $OH_{aq}$  into  $H_2O_{2aq}$  is shown in the lower panels. Notice that the location of the high concentration of OH at  $1\mu$ s becomes the location of the highest concentration of  $H_2O_{2aq}$ . Contours are on a 5-decade log-scale.

for the touching and not-touching cases beyond the change in energy deposition  $(8.0 \times 10^{11} \text{ cm}^{-3}, 2.2 \times 10^{12} \text{ cm}^{-3}$ , and  $8.5 \times 10^{12} \text{ cm}^{-3}$  for -10 kV, -15 kV and -20 kV), the absence of spreading of the plasma on the liquid in the not-touching case significantly reduces the inventory of OH available. The OH mutually reacts to form H<sub>2</sub>O<sub>2</sub>, reacts with nitrogen atoms to form NO, solvates into the water to form OH<sub>aq</sub>, or blows away within a few ms with the flow.

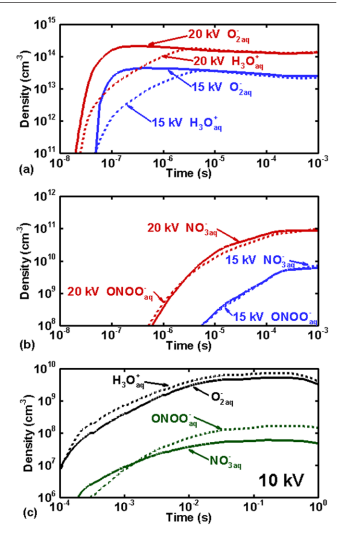
Photolysis of the water also produces  $OH_{aq}$ . The production of  $OH_{aq}$  by photolysis is less than 1% in the not-touching case and 10% in the touching case. The reason for more photolysis producing  $OH_{aq}$  in the touching case is that electron impact excitation produces  $N_2(b'\Pi_u)$  in the immediate vicinity of the surface of the water by the surface hugging plasma. VUV photons emitted by  $N_2(b'\Pi_u)$  are then able to reach the water prior to being absorbed in the humid air. By mutual reaction of  $OH_{aq}$ ,  $H_2O_{2aq}$  is produced which is also largely a terminal species in the absence of organic matter in the liquid. As shown in figure 6, the high concentrations of OH at 1 $\mu$ s resulting from



**Figure 7.**  $N_xO_y$  density for the  $-15 \,\text{kV}$  case shown at varying times after the discharge pulse in the gas and water layer. NO is the initiating molecule for producing  $N_xO_y$ . The comparatively low density of  $N_xO_y$  results from the effectiveness of the helium jet blocking access of NO to the water layer and the laminar flow that blows NO away. Contours are on a 4-decade log-scale.

spreading of the plasma through the evaporating water vapor produces a distribution of  $OH_{aq}$  having a maximum density of 2.2 × 10<sup>14</sup> cm<sup>-3</sup> or (3.7 × 10<sup>-7</sup> M) at a radius of 1 mm. The primary loss mechanism for  $OH_{aq}$  is the reaction forming  $H_2O_{2aq}$ ,  $OH_{aq} + OH_{aq} \rightarrow H_2O_{2aq}$ . The lifetime of  $OH_{aq}$  based on this maximum density and rate coefficient of 5.5 × 10<sup>9</sup> M<sup>-1</sup> s<sup>-1</sup> [14, 24] is about 0.25 ms. The resulting  $H_2O_{2aq}$  then diffuses through the water layer to the underlying tissue.

Reactive nitrogen species (RNS),  $N_xO_y$ , consist of the sum of NO, NO<sub>2</sub>, NO<sub>3</sub>,  $N_2O_3$ ,  $N_2O_4$  and  $N_2O_5$ . RNS are formed as the plasma plume reacts with the ambient air as shown in figure 7. Nitrogen oxide (NO) is initially formed primarily by reaction of N atoms with O<sub>2</sub>. The N atoms are formed dominantly by the electron impact dissociation of N<sub>2</sub> resulting from air diffusing into the plume. Since the He flow from the jet displaces air at the surface of the water and there is no N<sub>2</sub> injected through the tube, the RNS are primarily produced at the edge of the plasma jet where air diffuses into the plume. The precursor species to N<sub>x</sub>O<sub>y</sub>, NO, is largely formed at the boundary of the plasma where the density of the N<sub>2</sub> is largest and secondarily in the core of the plasma plume where the O



**Figure 8.** The time evolution of the liquid volume average densities of aqueous ions. (*a*)  $O_2^{-}_{aq}$  and  $H_3O_{aq}^+$ ; and (*b*)  $NO_{aq}^{-}_{aq}$  and  $ONOO_{aq}^{-}_{aq}$  for the touching -15 kV and -20 kV cases. (*c*) Densities for the -10 kV (not-touching) case.

atom density is the highest. The  $N_xO_y$  that are formed from the plasma discharge reacting with the ambient air react further react with OH and HO<sub>2</sub> in the ambient to form HNO<sub>2</sub>, HNO<sub>3</sub> and HOONO. The large densities of  $N_xO_y$  produced in, for example, DBDs accumulates over thousands of pulses and can reach  $3 \times 10^{17}$  cm<sup>-3</sup> in DBDs sustained in non-flowing air [25]. In the absence of vortices or other forms of recirculation in the gas phase, the NO and NO<sub>2</sub> that would normally be the starting point for forming higher  $N_xO_y$  in stagnant systems are blown out of the volume in plasma jets. The density of higher  $N_xO_y$  and acids HNO<sub>x</sub> are therefore small compared to DBDs, reaching a maximum density 1 mm above the liquid of only  $2 \times 10^{11}$  cm<sup>-3</sup> during the single pulse.

#### 3.2. Evolution of aqueous reactivity

The gas phase species in contact with the water then solvate to form aqueous species. The densities of aqueous ions averaged over the water layer as a function of time are shown in figure 8 for the -10 kV, -15 kV and -20 kV cases. Positive water ions are produced by charge exchange from gas phase

the negative ions.

positive ions striking the water and by photoionization. The  $H_2O_{aq}^+$  then quickly charge exchange with water to produce hydronium,  $H_3O_{aq}^+$  and hydroxyl radicals,  $OH_{aq}$ .  $H_3O_{aq}^+$  is a terminal species that dominates the positive ions and whose density is largely established by 1 ms after the discharge pulse. The average density of  $H_3O_{aq}^+$  in the water layer is  $2 \times 10^{14}$  cm<sup>-3</sup> and  $3 \times 10^{13}$  cm<sup>-3</sup> for the -20 kV and -15 kV cases. The aqueous anions are initially formed by solvated electrons dissociatively attaching with H<sub>2</sub>O<sub>aq</sub> to form OH<sup>-</sup>  $_{aq}$  or by attaching to the dissolved  $O_{2aq}$  to form  $O_2^{-}_{aq}$ . The  $OH_{aq}^{-}$  then combines with  $H_3O_{aq}^{+}$  or  $OH_{aq}$  to form  $OH_{aq}$  and  $O_{aq}^{-}$  respectively. The  $O_{aq}^{-}$  then combines with  $O_{2aq}$  to form the terminal negative ion  $O_3^-aq$ . In the absence of significant densities of  $N_x O_{yaq}$ , the long lived  $O_2^-_{aq}$  and  $O_3^-_{aq}$  dominate

As shown in figure 8 for -20 kV,  $O_2^-$  is quickly formed during the discharge pulse by attachment of solvated electrons to dissolved O<sub>2</sub> as the plasma touches and spreads on the surface of the water. The time for electron solvation is only tens of ps and the density of solvated electrons reaches its maximum value of 2.1  $\times$  10<sup>14</sup> cm<sup>-3</sup> within 200 ns. The density of H<sub>3</sub>O<sup>+</sup><sub>aq</sub> develops over a few  $\mu$ s since up to two charge exchange reactions are required for its formation-solvating ions reacting with water to form  $H_2O_{aq}^+$ , and then a second reaction with water to form  $H_3O_{aq}^+$ . There is also a contribution to  $H_3O_{aq}^+$  by hydrolysis of HNO<sub>xaq</sub>, however since the density of the acids is low, this is a minor contribution. The densities of  $O_2^{-}_{aq}$  and  $H_3O_{aq}^+$  for -15 kV show the same trends as for -20 kV. The initial rise in densities for  $-15 \,\text{kV}$  are delayed in time by about 20 ns compared to  $-20 \,\text{kV}$ , which is the difference in arrival times of the electron plumes at the surface of the water. The densities are also decreased by a factor of 5-6, a consequence of the shorter length of time that the surface ionization wave spreads on the water at the lower voltage.

The electron plume does not reach the water layer with a significant density for the -10 kV case and so there is no prompt formation of aqueous ions. As noted above, the electron-ion plume transitions to an ion-ion plume as recombination and attachment reduce the electron density. This recombining ionion plume then intersects with the water layer on up to a ms timescale, which is the time required for the plume containing the ion-ion plasma to cross the gap. Upon intersection with the water, the low density of gas phase ions  $(10^{10} \text{ cm}^{-3} \text{ at } 10 \mu \text{s})$ solvate into the water over several ms, producing average aqueous densities of  $>10^9$  cm<sup>-3</sup>. O<sub>2</sub><sup>-</sup><sub>aq</sub> and H<sub>3</sub>O<sup>+</sup><sub>aq</sub> are produced in a similar manner as described above for the higher voltages, evolving over 10s to 100s ms to produce an aqueous density of nearly 10<sup>10</sup> cm<sup>-3</sup>, a factor of 10<sup>4</sup> smaller than for the touching cases. This same, longer time solvation of the ion-ion plume also occurs for the higher voltages. However, the contribution of the ion-ion plasma plume to aqueous ion formation is simply smaller than the direct solvation of the electron-ion plasma in contact with the water.

The  $N_xO_v$  that reaches the water also solvates into the liquid, where N<sub>x</sub>O<sub>yaq</sub> eventually forms HNO<sub>2aq</sub>, HNO<sub>3aq</sub> and HOONO<sub>aq</sub> [14]. HNO<sub>2aq</sub> is a weak acid so only about 1% of its concentration hydrolyzes to  $H_3O_{aq}^+$  and  $NO_{2aq}^-$ . HNO<sub>3aq</sub> and HOONOaq are strong acids and nearly completely hydrolyze to S A Norberg et al

 $N_x O_{yaq}$  species are  $NO_3^-$  and  $ONOO^-$  aq. Since the majority of  $N_x O_y$  that solvates in the water is not formed in the spreading plasma on the surface, these molecules must convect and diffuse from more remote locations. This convection requires about 1 ms for the touching cases, resulting in the NO3-aq and  $ONOO_{aq}^{-}$  reaching a maximum density of  $10^{11}$  cm<sup>-3</sup> for -20 kV and  $10^{10}$  cm<sup>-3</sup> for -15 kV. The same process requires tens of ms in the not-touching cases producing a density of 10<sup>8</sup> cm<sup>-3</sup>. The dynamics of N<sub>x</sub>O<sub>y</sub> for DBDs in stagnant gases and in jets do significantly differ. In stagnant DBDs, N<sub>x</sub>O<sub>y</sub> accumulates in the gas phase, diffuses into the water and hydrolyzes. In jets, there are shorter residence times for the  $N_xO_y$  in the gas phase due to the convective flow and so less opportunity to accumulate to large densities. The result is less hydrolysis in the water and low densities of nitrates and nitrites in the liquid. For both types of plasmas sources, N<sub>x</sub>O<sub>vaq</sub> will accumulate in the liquid phase over successive discharge pulses.

The densities of neutral aqueous species averaged over the water layer as a function of time are shown in figure 9 for the -10kV, -15kV and -20kV cases. In the touching cases (-15 kV, -20 kV), the OH<sub>aq</sub> has a rapid rise during the discharge pulse (tens of ns) due to photolysis, augmented on the hundreds ns time scale by charge exchange reactions forming  $H_3O_{aq}^+$  and  $OH_{aq}$ . The maximum densities for  $OH_{aq}$  are  $5.8 \times 10^{14} \text{cm}^{-3}$  for -20 kV and  $7.4 \times 10^{13} \text{ cm}^{-3}$  for -15 kV. The majority of OH<sub>aq</sub> is expended in forming H<sub>2</sub>O<sub>2aq</sub>, whose maximum densities of  $2.7 \times 10^{14} \text{ cm}^{-3}$  for -20 kV and  $4.4 \times 10^{13} \text{ cm}^{-3}$  for -15 kV are reached in about 1 ms. The H<sub>aq</sub> that is produced by photolysis of H<sub>2</sub>O<sub>aq</sub> reacts with the dissolved  $O_{2aq}$  to form  $HO_{2aq}$ , which is also formed by  $O_{aq}$ reacting with OH<sub>aq</sub>. As the OH<sub>aq</sub> converts to H<sub>2</sub>O<sub>2aq</sub> over time in the water layer, the primary loss mechanisms for HO<sub>2aq</sub> become reactions with NO<sub>aq</sub> to form HNO<sub>3aq</sub> or hydrolysis to form  $H_3O^+_{aq}$  and  $O_2^-_{aq}$ . The density of  $HO_{2aq}$  rises during the discharge pulse due primarily to the source of H<sub>aq</sub> in photolysis reactions (6  $\times$  10<sup>14</sup> cm<sup>-3</sup> for -20 kV and 7  $\times$  10<sup>13</sup> cm<sup>-3</sup> for -15kV) and slowly declines on ms timescales as NO solvates to form NO<sub>aq</sub>. Since the source of O<sub>3aq</sub> is dominated by diffusion of O<sub>3</sub> from the gas phase, its density increases over many ms with maximum values of  $2.1 \times 10^{14} \text{ cm}^{-3}$ for -20 kV and  $5.6 \times 10^{13} \text{ cm}^{-3}$  for -15 kV.

The time evolution of OH<sub>aq</sub> for the not-touching case (-10kV) significantly differs from the touching cases. Lacking the photolysis and charge exchange reactions which produce OH<sub>aq</sub> on ns timescales as for touching cases, diffusion of OH into the water and dissociative excitation transfer to water molecules by N<sub>2</sub>(A) produce OH<sub>aq</sub> over hundreds of  $\mu$ s. Note that the rise of O<sub>3aq</sub> shadows that of OH<sub>aq</sub> as both require diffusion of neutral species from the gas phase for their formation in the liquid. The maximum densities of  $O_{3aq}$  and  $OH_{aq}$  are 6.4 × 10<sup>12</sup> cm<sup>-3</sup> and 2.8 × 10<sup>12</sup> cm<sup>-3</sup>. The OH<sub>aq</sub> converts to H<sub>2</sub>O<sub>2aq</sub> whereas O<sub>3aq</sub> is a terminal species. The density of  $OH_{aq}$  falls below that of  $H_2O_{2aq}$  within  $1\,\mu$ s, in agreement with Tresp *et al* [7]. The formation of the other major neutral aqueous species is less dependent on the plasma striking the surface than for ion species. An exception is  $HO_{2aq}$  whose formation benefits from the production of  $H_{aq}$ 

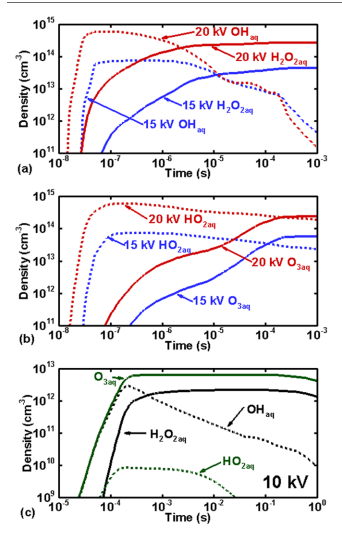
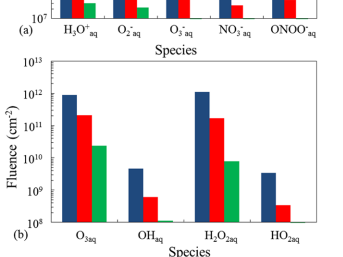


Figure 9. The time evolution of the liquid volume average densities of neutral aqueous species (a)  $H_2O_2aq$  and  $OH_{aq}$ ; and (b)  $O_3aq$  and  $HO_{2aq}$  for the touching -15 kV and -20 kV cases. (c) Densities for the  $-10 \,\text{kV}$  (not-touching) case.

during photolysis reactions which have at best a minor role in the not-touching case. As a result, HO<sub>2aq</sub> is disproportionately smaller for the -10kV case.

#### 3.3. Fluences to the tissue

The treatment of the underlying tissue ultimately depends on the fluence of reactive species that transport through the liquid layer. These fluences are shown in figure 10 after 10s following a single discharge pulse. The maximum fluences for  $H_3O_{aq}^+$  and  $O_{2aq}^-$  for -20 kV are  $3 \times 10^{11} \text{ cm}^{-2}$ . For the -15 kVtouching case, the fluences are reduced by a factor of 10. This reduction is largely a consequence of the shorter time the plasma spreads on the liquid for -15 kV. The production of aqueous ions by the ion-ion plume for  $-10 \,\text{kV}$  is smaller by a factor of  $10^4$ . The low rate of solvation of  $N_x O_y$  to form aqueous RNS is reflected in the small fluences of NO<sub>3<sup>-</sup>ag</sub> and ONOO<sup>-</sup>ag to the tissue, and which is only significant in the high voltage case. The neutral species are less affected by the touching or not-touching conditions. For example, the formation of  $O_{3aq}$ 



O<sub>3</sub>-aq

1012

1011

1010

109

 $10^{8}$ 

 $H_3O_{aq}^+$ 

Fluence (cm<sup>-2</sup>)

Figure 10. Fluences to the tissue beneath the water over 10 s for not-touching (-10kV) and touching (-15kV, -20kV) cases. (a) Ions and (b) neutral species.

results from the advective flux of O<sub>3</sub> from above the water layer, a process which is not particularly sensitive to touching or not-touching conditions. So the fluence of  $O_{3aq}$  is largely a function of energy deposition and decreases in proportion to the applied voltage (4 times more energy per pulse for -20kV compared to -10 kV). The fluence of  $H_2O_{2aq}$  to the tissue has a larger decrease with decreasing voltage than  $O_{3aq}$  due to the contribution of photolysis and charge exchange to formation of its precursor  $OH_{aq}$ . However this decrease in  $H_2O_{2aq}$  is small compared to the ions-a consequence of a large fraction of OH<sub>aq</sub> originating from diffusion of neutral species into the water. The fluence of  $HO_{2aq}$  for the not-touching case is not appreciable, as its formation depends on the production of H atoms by photolysis which is only significant for touching cases.

### Another factor which is sensitive to the plasma jet touching or not-touching the liquid is the delivery of electric fields to the tissue underlying the liquid. The maximum electric field at the surface of the tissue occurs at the end of the discharge

pulse for the -10 kV, not-touching, case and is 1 kV cm<sup>-1</sup>. For the touching cases, the maximum electric field at the surface of the tissue occurs within 5-10ns of the ionization wave striking the water layer and reaches a value of 14 kV cm<sup>-1</sup> for the -15 kV case and  $17 \text{ kV} \text{ cm}^{-1}$  for the -20 kV case. From

3.4. Electric fields

-20 kV

-15 kV

-10 kV

ONOO<sup>-</sup>aq

shortly after impact of the ionization wave until the pulse terminates, the magnitude of the electric field decreases by only a few  $kV \text{ cm}^{-1}$ . Once the voltage is removed, the field quickly dissipates as the charge density on the surface of the water decreases. These values of electric field for a single pulse, up to  $20 kV \text{ cm}^{-1}$  for tens of ns, are near the threshold for producing electroporation [26]. The threshold electric field for electroporation decreases with increasing number of pulses, and so the potential for producing electroporation with touching plumes may be significant at high PRF.

#### 4. Concluding remarks

One of the major distinctions in producing active aqueous species in water layers over tissues by plasma jets is whether the visible electron-ion plasma plume touches the surface of the water. There are at least two major modes for delivery of active species to the water layer. The first can dominate in touching cases-the spreading of plasma over the liquid surface. This immediate proximity of the plasma to the liquid enables photolysis, direct charge exchange reactions with H<sub>2</sub>O<sub>a0</sub>, and direct solvation of electrons. The longer the dwell time of the plasma on the surface of the liquid, the more aqueous ions and photolysis products are formed over a larger area. Species such as  $HO_{2aq}$  that trace their origins to these reactions are also produced in larger numbers. Not-touching plasmas do receive fluxes of gas phase ions in the form of an ion-ion plume, however these fluxes can be orders of magnitude smaller than in the touching cases.

The second mode of delivery is convection and diffusion of neutral species to the liquid which are formed remotely from the surface. The formation of aqueous reactivity by neutral species is much less sensitive to touching or not-touching conditions, as demonstrated by the commensurate, voltage scaled fluences of  $O_{3aq}$  to the tissue discussed here. The delivery of these neutral species to the water is more sensitive to fluid dynamics than for the ions. Vortices, turbulence and other forms of recirculation that extend the residence time of neutral species at the surface of the water increases the likelihood of their solvation. This is particularly the case for N<sub>x</sub>O<sub>y</sub> species that typically accumulate over many discharge pulses in DBDs sustained in non-flowing gas. For strictly laminar, nonvortexed flows,  $N_x O_v$  must diffuse through the He core of the jet to reach the surface, and is more likely to be blown away before solvating in significant amounts.

#### Acknowledgements

This work was supported by the Department of Energy Office of Fusion Energy Science (DE-SC0001319) and the National Science Foundation (CHE-1124724).

#### References

[1] Laroussi M, Kong M, Morfill G and Stolz W 2012 Plasma Medicine: Applications of Low-Temperature Gas Plasmas *in Medicine and Biology* (Cambridge: Cambridge University) pp 239–60

- [2] Kong M G, Kroesen G, Morfill G, Nosenko T, Shimizu T, van Dijk J and Zimmerman J L 2009 Plasma medicine: an introductory review *New J. Phys.* 11 115012
- [3] Laroussi M 2009 Low-temperature plasmas for medicine? *Trans. Plasma Sci.* 37 714
- [4] Rhoades R A and Bell D R 2012 Medical Physiology: Principles for Clinical Medicine 4th edn (Baltimore: Williams & Wilkins) pp. 167–77
- [5] van Gils C A J, Hofmann S, Boekema B K H L, Brandenburg R and Bruggeman P J 2013 Mechanisms of bacterial inactivation in the liquid phase induced by a remote RF cold atmospheric pressure plasma jet *J. Phys. D: Appl. Phys.* 46 175203
- [6] Lukes P, Dolezalova E, Sisrova I and Clupek M 2014Aqueousphase chemistry and bactericidal effects from an air discharge plasma in contact with water: evidence for the formation of peroxynitrite through a pseudo-second-order post-discharge reaction of H<sub>2</sub>O<sub>2</sub> and HNO<sub>2</sub> *Plasma Sources Sci. Technol.* 23 015019
- [7] Tresp H, Hammer M U, Winter J, Weltmann K-D and Reuter S 2013 Quantitative detection of plasma-generated radicals in liquids by electron paramagnetic resonance spectroscopy *J. Phys. D: Appl. Phys.* 46 435401
- [8] Lu X, Naidis G V, Laroussi M and Ostrikov K 2014 Guided ionization waves: theory and experiments *Phys. Rep.* 540 123
- [9] Reuter S, Tresp H, Wende K, Hammer M, Winter J, Masur K, Schmidt-Bleker A and Weltmann K-D 2012 From RONS to ROS: tailoring plasma jet treatment of skin cells *Trans. Plasma Sci.* 40 2986
- [10] Reuter S, Winter J, Schmidt-Bleker A, Tresp H, Hammer M and Weltmann K-D 2012 Controlling the ambient air affected reactive species composition in the effluent of an argon plasma jet *Trans. Plasma Sci.* 40 2788
- [11] Bruggeman P 2014 private communication
- [12] Lay B, Moss R, Rauf S and Kushner M 2003 Breakdown processes in metal halide lamps *Plasma Sources Sci. Technol.* 12 8
- [13] Xiong Z and Kushner M J 2010 Surface corona-bar discharges for production of pre-ionizing UV light for pulsed highpressure plasmas J. Phys. D: Appl. Phys. 43 505204
- [14] Tian W and Kushner M J 2014 Atmospheric pressure dielectric barrier discharges interacting with liquid J. Phys. D: Appl. Phys. 47 165201
- [15] Bhoj A N and Kushner M J 2007 Continuous processing of polymers in repetitively pulsed atmospheric pressure discharges with moving surfaces and gas flow J. Phys. D 40 6953
- [16] Walsh J and Kong M 2008 Contrasting characteristics of linear-field and cross-field atmospheric plasma jets *Appl. Phys. Lett.* **93** 111501
- [17] Lu X, Ziang J, Xiong Q, Targ Z, Hu X and Pan Y 2008 An 11cm long atmospheric pressure cold plasma plume for applications is plasma medicine *Appl. Phys. Lett.* **92** 081502
- [18] Maletic D, Puac N, Lazovic S, Malovic G, Gans T, Schulz-von der Gathen V andLj Petovic Z 2012 Detection of atomic oxygen and nitrogen created in a radio-frequencydriven micro-scale atmospheric pressure plasma jet using mass spectroscopy *Plasma Phys. Control. Fusion* 54 124046
- [19] Robert E, Sarron V, Darny T, Ries D, Dozias S, Fontane J, Joly L and Pouvesle J-M 2014 Rare gas flow structuration in plasma jet experiments *Plasma Sources Sci. Technol.* 23 012003
- [20] Bogey C and Bailly C 2010 Direct and large-eddy simulation VII Proceedings of the Seventh International ERCOFTAC Workshop on Direct and Large Scale Eddy Simulation VII (Trieste, Italy) ed V Armenio et al (Berlin: Springer)

- [21] Karakas E, Akman M A and Laroussi M 2012 The evolution of atmospheric-pressure low-temperature plasma jets: jet current measurements *Plasma Sources Sci. Technol.* 21 034016
- [22] Jiang N, Ji A and Cao Z 2010 Atmospheric pressure plasma jets beyond ground electrode as charge overflow in a dielectric barrier discharge setup J. Appl. Phys. 108 033302
- [23] Hubner S, Hofmann S, van Veldhuizen E M and Bruggeman P J 2013 Electron densities and energies of a guided argon streamer in argon and air environments *Plasma Sources Sci. Technol.* 22 065011
- [24] Madden K P and Mezyk S P 2011 A critical review of aqueous solution reaction rate constants for hydrogen atoms J. Phys. Chem. Ref. Data 40 023103
- [25] Sakiyama Y, Graves D, Change H-W, Shimizu T and Morfill G E 2012 Plasma Chemistry model of surface micro-discharge in humid air and dynamics of reactive neutral species *J. Phys.* D 45 425201
- [26] Beebe S J, Blackmore P F, White J, Joshi R P and Schoenbach K H 2004 Nanosecond pulsed electric fields modulate cell function through intracellular signal transduction mechanisms *Physiol. Meas.* 25 1077