Corona discharges with water electrospray for *Escherichia coli* biofilm eradication on a surface

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**A B S T R A C T**

Low-temperature plasma (cold), a new method for the decontamination of surfaces, can be an advantageous alternative to the traditional chemical methods, autoclave or dry heat. Positive and negative corona discharges in air were tested for the eradication of 48-h *Escherichia coli* biofilms grown on glass slides. The biofilms were treated by cold corona discharge plasma for various exposure times. Water electrospray from the high voltage electrode was applied in some experiments. Thermostatic cultivation of the biofilm, and confocal laser scanning microscopy (CLSM) of the biofilm stained with fluorescent dyes were used for biocidal efficiency quantification. Up to 5 log_{10} reduction of bacterial concentration in the biofilm was measured by thermostatic cultivation after exposure to both corona discharges for 15 min. This decontamination efficiency was significantly enhanced by simultaneous water electrospray through the plasma. CLSM showed that the live/dead ratio after treatment remained almost constant inside the biofilm; only cells on the top layers of the biofilm were affected. DAPI fluorescence showed that biofilm thickness was reduced by about 1/3 upon exposure to the corona discharges with electrospray for 15 min. The biofilm biomass loss by about 2/3 was confirmed by crystal violet assay.

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1. Introduction

Bacterial contamination of surfaces is a common problem in hospitals, the food industry, water distribution systems, etc. Nosocomial infections (NI), or so-called hospital acquired infections are responsible for infecting 2–11% of the patients admitted to the hospital in the European Union (EU) and approximately 37,000 deaths/year are caused directly by NI in the EU 27 and contribute to an additional 111,000 deaths/year. *Escherichia coli* is one of the most frequently isolated strains from all infection sites in health care facilities [1]. Bacteria on surfaces exist predominantly in the form of biofilms. Microbial biofilms are populations of microorganisms concentrated at an interface (usually solid-liquid) and encased in a hydrated matrix of exopolymers such as EPS, polysaccharides, and proteins that are produced by the resident microorganisms [2]. EPS protects cells from the outer environment and facilitates cell-to-cell communication (quorum sensing) [3]. Quorum sensing is required for biofilm differentiation [4], it coordinates gene expression and regulates a diverse array of physiological activities [5]. Bacteria in the biofilm are protected from harsh conditions (high temperature, low pH, ultraviolet radiation, dehydration, etc.), and therefore are more resistant than their planktonic (suspended cells) counterparts [3]. In order to avoid using toxic chemicals or high concentrations of antibiotics to achieve the desired decontamination efficiency in biofilms, a search for new alternative methods of decontamination is required; one of these is low-temperature plasma.

A plasma is defined as a macroscopically neutral ionized gas. Most plasmas found in nature exist at high-temperature and are in thermodynamic equilibrium. In this article, the focus will be on non-thermal plasma or so-called non-equilibrium or low-temperature (cold) plasma. In cold plasma, the temperature of electrons is high, while the temperature of heavy particles (atoms, molecules, and ions) remains close to ambient. This type of plasma can be generated at atmospheric pressure [6,7]. Low-temperature plasmas studied under laboratory conditions are typically generated by electrical discharges in helium [8] or argon [9,10] with admixtures of reactive gases such as oxygen or water vapor [11,12], or they can be directly generated in air [13–16]. In cold non-thermal atmospheric pressure plasma sources, the major active biocidal agents are reactive neutral species (reactive oxygen and nitrogen species, when plasma is produced in air), UV–Vis radiation, and charged species (electrons and ions) along with the corresponding electromagnetic fields [17].

Non-thermal plasma at atmospheric pressure is well-suited for decontamination of thermally sensitive surfaces because the bulk
temperature remains close to ambient. Its bactericidal effects have been previously tested on a wide range of bacterial species - planktonic bacteria [18-23], spores [21,24-28] or bacterial biofilms [14,29-40]. Thanks to the ease of use of plasma on target surfaces, the first plasma medicine applications appeared in dermatology for wound healing [10,41] and blood coagulation [42], in dentistry [43,44] for root canal disinfection [45], and in dental aesthetics for tooth whitening [46].

To increase the production and amounts of the reactive neutral species, water can be added to the discharge, in the form of water vapor (as mentioned before) or as fine liquid droplets electrosprayed from the high-voltage electrode. The electrospray effect emerges when the liquid flowing out from a capillary is exposed to high electrical potential. A “Taylor cone” is produced and the droplets of liquid emerging from its tip being charged are thus accelerated by the electric field [47-50]. The electrosprayed water brings more complexity to the discharge chemistry and its interaction with bacteria. The effect of water electrospray combined with corona discharges has been previously studied in our group [47,48,51] and applied to polymer surfaces contaminated by biofilms and spores [13,52] and to water disinfection [15,53]. In this paper, we investigate the impact of water electrospray and polarity of the air corona discharge on 48-h Escherichia coli biofilm. The biofilm decontamination, thickness reduction, and biomass loss are evaluated by various microbiological and fluorescent microscopy methods.

2. Materials and methods

Non-thermal (cold) plasma generated by both positive corona (PC) and negative corona (NC) discharges in air were applied for the eradication of Escherichia coli biofilm formed on glass surfaces (cover slides). The effect of the water electrospray combined with the discharge on its bactericidal efficiency was also tested.

2.1. Experimental set-up and discharges

Corona discharges in atmospheric pressure air were generated in an experimental set-up consisting of a needle-to-plane electrode system placed in a discharge chamber in open air. The needle electrode was a sharp or a clipped hypodermic syringe needle connected to a DC high-voltage (HV) power supply. Opposite the needle HV electrode, a copper plate was grounded through a 50 kΩ resistor. The treated biofilm samples on glass cover slides were placed on the grounded electrode, 5 mm from the HV electrode. Some experiments were performed with sterile distilled water electrosprayed onto the sample through a hollow clipped HV needle electrode; this was in turn supplied to the needle by pumping with a NE-300 SyringePump (Fig. 1). The electrical characteristics of the discharges were measured as follows: the applied voltage with a Tektronix P6015A HV probe connected to the needle electrode, and the discharge current by measuring the voltage across the 50 kΩ grounded resistor. Both probes were connected through coaxial cables to a Tektronix TDS 2024 digital oscilloscope for signal recording and storage.

Corona discharges of both polarities were operated in the configuration detailed above. Positive corona (PC) was supplied with a voltage up to +9 kV and formed streamers with frequencies ranging from 10 to 20 kHz and maximum current pulse amplitudes up to 10 mA (corresponding to a mean input power P ≈ 100 mW). With water electrospray the pulse frequency was lowered to 10 kHz, although the current amplitude was slightly increased but remained under 50 mA (P ≈ 200 mW). Negative corona (NC) was supplied with a minimum voltage of −9 kV and current pulses with frequencies ranging from 0.5 to 2 MHz and amplitudes of −1 mA were observed (P ≈ 300 mW). With water electrospray, the pulse frequency was lowered to 200–500 kHz and the current amplitude increased to −10 mA (P ≈ 400 mW). More details on the discharge experimental conditions can be found in [13].

2.2. Bacterial samples

Escherichia coli strain BW25113 F− [BD792 derivative, rnb, DElacZ4787, HsdR514, DE(araBAD)567, DE(araBAD)568, rph−1] was used to form a biofilm on glass cover slides (2 cm × 2 cm × 15 μm) on the bottom of a homemade 6-well plate for 48 h at 30 °C. A stationary phase culture grown in Miller’s modified Luria broth (LB) (37 °C) was diluted 1:100 in buffered (pH 7.4 Phosphate buffered solution) M63 medium (AMRESCO) supplemented with 0.1% casamino acids and 1 g/L glucose. One milliliter of this suspension was placed into each well of the 6-well plate. The samples were incubated without agitation at 30 °C for 48 h with media refreshment after 24 h.

2.3. Plasma treatment and sample post-treatment

The biofilms on glass cover slides were taken out from the 6-well plates after 48 h. The excess liquid was carefully removed, the samples were dried for up to 20 min at 35 °C, and then placed onto the grounded electrode inside the discharge chamber and treated with plasma. The

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**Fig. 1.** Photographs of the DC corona discharges in air with a hypodermic injection needle as HV electrode (clipped for water electrospray b, d) and a glass cover slide on a copper grounded electrode: a) positive corona, b) positive corona with water electrospray, c) negative corona, d) negative corona with water electrospray (4 s exposure time).
control samples were dried using the same procedure but not treated with plasma.

Cultivability was determined by repetitive rinsing of the biofilm with 5 mL of 0.85% NaCl saline solution and scraping with another sterile cover glass. The recovered bacterial cells in the solution were vortexed, serially diluted and spread over LB agar in petri dishes and incubated at 37 °C overnight, then the bacterial colony forming units (CFUs) were counted.

For imaging using confocal laser scanning microscopy (CLSM), the treated biofilms were stained with three fluorescent dyes: 1 μL Syto9 (5 mmol·L\(^{-1}\), Life Technologies), 1.5 μL Propidium iodide (PI, 20 mmol·L\(^{-1}\), Cayman chemicals) and 5 μL DAPI (4′,6-diamidino-2-phenylindole, 2 mg/mL, Cayman chemicals) in phosphate buffered saline (PBS, pH 7.4), and incubated for 25 min in the dark.

CLSM images were acquired with an OLYMPUS IX81 inverted confocal laser microscope. Z-stack acquisitions were performed at 16 bits, with dimensions of 640 × 640 × 1 pixels (pixel size 0.33 × 0.33 × 0.4 μm), the dimension of the final z-stack was 211.2 × 211.2 × (0.4 × N) μm, where N is the number of z optical slices taken. Observed excitation and emission wavelengths (filters) as follows: Propidium iodide: excitation 559 nm/emission 575–675 nm, Syto9: 473 nm/490–540 nm, DAPI: 405 nm/430–455 nm. Image stacks (.oif or .oib format) were analyzed using Icy 1.6.1.1v [54] from which the sum of intensity in each channel (red—PI, green—Syto9, blue—DAPI) of each optical slice in the stack and three dimensional projections of stacks were obtained.

E. coli BW 25113 F+ in planktonic form was used to obtain the calibration curve for live/dead ratios (according to the supplier manual). Briefly, bacteria in late-log phase were centrifuged to remove nutrient broth and split into two parts: one part with live bacteria diluted in 0.85% NaCl saline solution, and the other part in 70% 2-propanol to kill bacteria. After incubation and washing, the live and dead bacteria were mixed together in various ratios, then stained, and the fluorescence was measured using CLSM. The calibration curve was constructed from the ratios of green (Syto9) and red (PI) fluorescence for the known bacteria live/dead ratios (Fig. 2). A real live/dead ratio can be estimated from this curve.

2.4. Biofilm biomass evaluation

Biofilm biomass was evaluated using crystal violet (CV) staining following the established microbiology protocol [55]. The control and plasma-treated biofilms were reseeded in the 6-well plate after treatment and 200 μL of 0.1% crystal violet was introduced into each well. After 10 min incubation, the CV solution was carefully removed by pipetting and biofilms were rinsed with deionized water until the waste liquid was clear. The 6-well plate was dried upside-down overnight at room temperature. When fully dried, 200 μL of 33% acetic acid was added into each well for 15 min to solubilize the CV, then recovered and diluted 1/10 in deionized water and the absorbance was measured at 550 nm.

3. Results

Results of cold plasma application to the biofilm evaluated by thermostatic cultivation and the analysis of biofilm structural changes by CLSM and CV assay are presented here.

3.1. Thermostatic cultivation

The reduction in bacterial colony forming units after plasma treatment with variable exposure time (2.5, 5, 10, and 15 min) is presented in Fig. 3. The initial bacterial concentration in the biofilm was 3.11 ± 6.59 × 10^6 CFU (colony forming units) per mL (the biofilms were disrupted and resuspended in 5 mL of NaCl solution). In both corona polarities, the decontamination efficiency increased with exposure time and reached up to 4.76 and 4.99 log_{10} reduction for positive and negative corona within 15 min, respectively. No significant differences between positive and negative corona were found, except for 5 min exposure time (p = 0.043, Mann-Whitney test).

To evaluate the efficiency of water electrospray through the discharge we present two exposure times: 2.5 and 15 min (Fig. 4). For 2.5 min exposure time, the electrospray increased the efficiency from 1.59 to 2.18 log_{10} for PC and 1.73 to 3.41 log_{10} for NC. For negative corona this change was found significant (p < 0.01, Mann-Whitney test). For the 15 min exposure time, the difference in efficiency between the plasma only treatment and the plasma with electrospray treatment was found to be smaller, but significant (PC, p = 0.048, Mann-Whitney test) or marginally significant (NC, p = 0.074, Mann-Whitney test). Decontamination efficiency increased from 4.76 to 5.28 log_{10} and from 4.99 to 5.4 log_{10} for PC and NC, respectively, when the water electrospray was used.

Fig. 2. Calibration curve for bacteria live/dead ratio (ratio of the fluorescence of Syto9 and PI) inside the biofilm measured for the known percentage of live planktonic bacteria (6–10 repeats for each point, mean ± 95% CI).

Fig. 3. Reduction of bacterial concentration in the biofilm on the cover glass by cold plasma treatment of the corona discharges (medians with IQR – interquartile range, 5–6 independent repetitions, * significant difference between polarities with a probability of error <0.05).
3.2. Confocal laser scanning microscopy

Images from CLSM contain information on the spatial distribution of the fluorescence from three different fluorescent dyes in the biofilm. DAPI binds preferably to dsDNA and its blue fluorescence is proportional to the amount of the present DNA. In our case DAPI was used to stain all cells in the biofilm. Red PI also stains DNA but does not penetrate through intact cell membranes and is therefore used to stain dead cells, or cells with damaged membranes. Syto9 is a green DNA dye which can penetrate inside all cells. Since PI has a stronger affinity to DNA than Syto9, when both are present in a cell, Syto9 is displaced from DNA by PI and cells are stained red (according to the protocol — L7012 LIVE/DEAD® BacLight™ Bacterial Viability Kit, Molecular Probes).

The fluorescence was acquired from four examined spots on the biofilm: the center and three random places around the center. Examples of reconstructed three-dimensional examined spots (211.2 × 211.2 μm) of the biofilm (the entire biofilm is a ∅ 12 mm disc) are presented in Fig. 5. From this representation we can see that the plasma treated biofilms contain more dead cells (red) on the top than the controls. The structure of the biofilm also changed with plasma treatment: the biofilm seems denser with more compact cellular structure. After plasma treatment with the water electrospray, the biofilm became thinner and patchy.

To quantify the effect of the plasma treatment on the biofilm, the live/dead ratio (Syto9/PI) was calculated. The sums of the red and green fluorescent intensities were measured in each optical slice of the biofilm. The relative values of fluorescence intensity ratios, corresponding to the real live/dead bacteria ratio obtained from the calibration curve (Fig. 2), were calculated and plotted along the depth position z in the biofilm (from the bottom z = 0 to the top of the biofilm — Fig. 6 — red). In the control samples (Fig. 6(a)), the live/dead ratio remained constant at 0.9 ± 0.3 in the bulk of the biofilm (from z = 0 to 17 μm), and increased up to 2 ± 1.5 toward the biofilm surface (z = 35 μm). After a 15 min plasma treatment (with or without electrospray), the mean live/dead ratio remained constant (around 0.7–0.8) for both polarities in the biofilm bulk, and the topmost layers of cells (with the highest live/dead ratio in the control) were lost. This corresponds to the reduction of biofilm thickness which decreased from 35 μm (control) down to 23 μm by PC with water electrospray, and down to 16 μm without water electrospray. For NC, the biofilm thickness was reduced from 35 μm (control) down to 13 μm with water electrospray, and down to 18 μm without water electrospray.

The weak influence of the plasma treatment on the live/dead bacteria ratio in the bulk of the biofilm may indicate that the plasma (active species and radiation) does not penetrate sufficiently inside the biofilm. This can be caused by cellular debris and EPS, which protect the deeper layers of bacteria in the biofilm. The values of live/dead ratio calculated from the total fluorescence of Syto9 and PI (not layer by layer but integrated through all layers) measured on all biofilm samples are presented in Fig. 7. These results from all experiments summed together (unlike one specific experiment shown in Fig. 6) showed a slight decrease in the live/dead ratio with plasma treatment that is only significant for PC 15 min exposure time with water electrospray. The live/dead ratio decreased from 0.91 ± 0.02 (control) to 0.61 ± 0.03 in PC, and did not change for NC, at 15 min exposure time and either with or without water electrospray.

The same visualization as for Syto9 and PI (live/dead ratio) was applied for DAPI, for which the sum of the blue fluorescence in the optical slice depends on the depth position in the biofilm (Fig. 6 — blue). Using the DAPI staining technique, in the control samples (Fig. 6a) the biofilm reached up to 30 μm in thickness. The obtained results with DAPI staining confirm the trend in the thickness reduction (biofilm shrinking) observed with the live/dead ratio technique. Again, the stronger effect was obtained for NC with electrospray.

To better quantify this loss of the biofilm thickness we chose to define the biofilm beginning and end as the slice z-positions with 10% fluorescence compared to the maximal fluorescence observed for all the
The difference between z position of the end and the beginning was established as the biofilm thickness. The values of the biofilm thickness were calculated for all z-stacks in all biofilms and the result mean values are presented in Fig. 8. There is a trend of reduction of the biofilm thickness with the rising exposure time and it is enhanced by adding the water electrospray to the discharge. The only significant difference was found for NC 15 min with electrospray in comparison with the control sample (p < 0.05, ANOVA test — multiple comparisons).

3.3. Biomass evaluation

The previous results demonstrate the reduction of the biofilm thickness upon corona discharge treatment. However, the observed biofilm thickness reduction was not very strong; we suppose that disrupted biofilm layers with dead cells still remained on the top of the intact biofilm. By alleviation (wash-out) of the detached bacteria we can observe the true loss of the biofilm biomass, which could have been underestimated by the presence of these detached but still present cells. The biofilm was stained with crystal violet (CV), incubated and the excess dye was rinsed together with the detached bacteria. The absorbance of solubilized CV was then measured. Considering the dispersion in biofilm (controls) thickness over all experiments, the biomass in plasma treated samples was calculated as the percentage of the control sample absorbance (control sample biomass is equal to 100%). All experiments shown in Fig. 9 were done for 15 min exposure time to the...
corona discharge. 46.4% and 33.7% of the biomass was preserved for PC and NC, respectively, i.e., remained attached to the surface after exposure to the discharges. When the water electrospray was added to the discharge, stronger bacteria detachment occurred and only 36.5% and 29.5% of the biomass remained attached to the cover glass in PC and NC treatment, respectively. All these correspond to significant reductions in comparison with the control samples (ANOVA multiple comparisons $p < 0.05$, for NC with electrospray $p < 0.01$).

4. Discussion

In the previous sections, two methods for biofilm viability evaluation were presented: a mechanical disruption of the biofilm followed by a thermostatic cultivation (CFU counts) of bacteria in the resulting solution, and a fluorescent staining and analyzing of the biofilm, layer by layer using the CLMS technique. The reduction of the biofilm thickness and the amount of the attached biomass were investigated by fluorescent DAPI staining, and by colorimetric crystal violet biomass assay. These measurements provided different results but all of them account for different effects of the plasma treatment of biofilms that are discussed in the following sections.

4.1. Biocidal efficiency

Thermostatic cultivation (CFU count method) showed a high biocidal efficiency for both polarities of the corona discharge treatment: almost 5 log$_{10}$ reduction of the bacterial population was achieved for a 15 min treatment time. Fig. 3 shows a slightly higher (not statistically significant) decontamination efficiency of NC. This was probably caused by higher discharge input power of NC (P $\approx$ 300 mW) in comparison with PC (P $\approx$ 100 mW). The only exposure time where PC was more efficient than NC was 10 min; in this case the mean power output of NC was lower than usual at 230 mW, but remained unchanged for PC. Thus, it can be concluded that PC was more energetically efficient than NC, since lower energy deposition of PC provided the same or even higher biocidal efficiency than NC. In PC, positive streamers propagate onto the biofilm surface, so producing in-situ active species (O atoms, OH radicals, and ions) and UV emission (UV-B). Similar neutral active species are also produced by NC, but only in the vicinity of the needle tip (not directly on the biofilm surface); the transport of these short-lived species to the cover glass placed on the plane electrode is supported by the ionic wind, but lower densities can be assumed to be present at the biofilm/gas interface. In addition, relatively stable species such as ozone and nitrogen oxides are produced in both polarities. These species can also contribute to the biofilm treatment.

By adding the water electrospray to the discharge, the decontamination efficiency increased up to 5.5 log$_{10}$ reduction. In this case, NC was slightly more efficient than PC (but not statistically significant). Energy deposited by the discharge on the sample increased for both polarities in this case, but was twice higher for NC (P $\approx$ 400 mW) than for PC (P $\approx$ 200 mW). In the discharge with water electrospray the tip of a hypodermic injection needle was clipped to enable the spray through the discharge. This resulted in the formation of two sharp points from which corona discharge was developed (Fig. 1b, d). Such a configuration can influence a larger surface of the treated sample, which is visible even from the photographs, and therefore caused higher decontamination efficiency in the experiments with water electrospray. There was also an increase of the power when the water was electrosprayed onto the sample, because a water layer was progressively build up on the cover glass and shortened the distance between the HV and the grounded electrode. Our present results do not clearly show whether the decontamination efficiency increase comes from the higher energy deposit or from other mechanisms such as: increase of the OH radical formation, re-hydration of the biofilm enhancing penetration of active species in its bulk, or others.

CLSM results showed that live/dead ratio development through the biofilms (Fig. 6) remains almost constant in the bulk of the biofilm after plasma treatment. The bacteria in the top layers are killed and their cell debris along with the EPS protect the bottom layers of cells by shielding them from direct contact with the plasma (PC). The overall live/dead ratio in biofilms (Fig. 7) confirmed that the plasma effect on the biofilm remained spatially superficial. These results contradict those obtained by the thermostatic cultivation. One hypothesis for explaining this contradiction are possible changes in bacterial metabolism after plasma treatment which might induce the viable but nonculturable state (VBNC), as previously hypothesized by Joaquín et al., 2009 [32]. This change can be induced by oxidative stress, or desiccation from the plasma source [35], visible light, starvation, osmotic stress [56] and other adverse conditions. The cytoplasmic membrane of bacteria in VBNC state may remain intact, therefore the PI fluorescent dye would not penetrate inside the cells and they may appear live in CLSM. On the other hand, VBNC bacteria in dormancy state will not proliferate and divide and thus will not form colonies on agar plates, which might cause the apparent high decontamination efficiency evaluated from CFU plate count. This has been previously reported: plasma treated bacterial cells in a biofilm [32,35] or a planktonic form [57,58] seem to be dead based on the culturability test, although the fluorescence and metabolic experiments showed that cells were intact and had a functional respiratory system. Therefore these bacteria could be still viable and may preserve their virulence [39].

4.2. Biofilm thickness and biomass reduction

The measurement of the biofilm thickness by DAPI fluorescent staining (Fig. 8) showed a significant reduction for a biofilm treated 15 min by the plasma with electrospray. However this decrease in the thickness was smaller than expected and did not agree well with the CV assay that showed almost 50% biomass loss (Fig. 9). One possible explanation might be that when bacteria are subjected to the stress they can become smaller [59] and more resistant to adverse environmental conditions [60]. The plasma treatment can also affect the EPS of the biofilm, the polymers surrounding the bacterial cells. As was previously reported in Vandervoot and Brelles-Marino 2014 [39], after longer exposure to the plasma the extracellular polymeric substance is removed or at least reduced. Since crystal violet only detects the attached biomass, the possibility that the observed biomass loss results from the detached biomass cannot be neglected. The plasma treatment impacted the
stability of the biofilm by reducing the EPS and the cell adhesion [35,39]. Since the biofilm was rinsed during the CV assay, loosely attached and detached bacteria were probably washed away. Rinsing was normally not applied for DAPI visualization, therefore the DAPI biofilm thickness measurements include the detached bacteria and debris in the plasma disrupted biofilm. For this reason, we performed an additional experiment with rinsing of the biofilm after the corona treatment. Both control and plasma treated samples (15 min, electrospray) were rinsed and compared with unrinsed samples (Fig. 10) by CLSM live/dead staining. Control samples remained almost intact (visible in Fig. 10d, side and bottom frames showing the depth profile of the fluorescent signal through the biofilm in corresponding x and y axes), while plasma treated samples were reduced almost to a monolayer of bacteria attached to the glass cover slide. This result confirms that corona discharges were able to disrupt the biofilm structure, which can be then easily detached by washing, while leaving only a monolayer of live single or clustered bacteria attached to the surface of the cover glass. We can assume that by repeating the procedure “plasma treatment + rinsing” we can completely decontaminate and clean the cover glass surfaces.

5. Conclusion

Low-temperature (cold) atmospheric pressure plasmas represent a new promising method for surface decontamination, alternative to the traditional chemical and heat treatment methods (autoclave, dry heat, etc.). The decontamination effects of the positive and negative corona discharges in air, examples of low-temperature plasmas, were tested on Escherichia coli biofilm grown for 48 h on glass slides. The biofilms were treated by the corona discharge plasma for various exposure times from 2.5 to 15 min. Additionally, water electrospray from the HV electrode onto the biofilm samples was used in some experiments. To quantify the biocidal efficiency of the discharges, different methods were used: thermostatic cultivation of bacteria from the biofilm scraped and disrupted in the solution and CFU counting, or confocal laser scanning microscopy (CLSM) of the biofilm stained with fluorescent dyes (Syto9, propidium iodide and DAPI). Bacteria live/dead fluorescence ratio vertical profiles were visualized through the biofilm. The fluorescence intensity of DAPI was used to evaluate the biofilm thickness loss, and the biofilm biomass loss was measured by the Crystal Violet assay.

Thermostatic cultivation of the bacteria from the biofilm showed 4.8 and 5 log10 reductions observed after 15 min exposure to positive and negative corona discharge, respectively. This decontamination efficiency was significantly enhanced by water electrospray onto the samples to 5.3 log10 for positive and to 5.4 log10 reduction for negative corona. CLSM showed that the live/dead ratio remained constant in the bulk of the biofilm after treatment, only cells on the top layers were affected by the discharge. Although the live/dead ratio did not decrease, the cells were destroyed and missing. The DAPI staining showed that the biofilm was thinner after plasma treatment (negative corona with electrospray for 15 min), with its thickness decreased from 18.1 μm in the control samples to 12.3 μm after treatment. A substantial biofilm biomass loss was confirmed by the Crystal Violet assay. Additionally, the live/dead staining of the plasma treatment combined with rinsing after the plasma exposure confirmed the substantial reduction of biofilm thickness, which indicates an interesting possibility for potential future applications.

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