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Effects of Nonequilibrium Atmospheric Pressure Plasmas on the Heterotrophic Pathways of Bacteria and on their Cell Morphology

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Effects of nonequilibrium atmospheric pressure plasmas on the heterotrophic pathways of bacteria and on their cell morphology

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To date, most research on the interaction of nonequilibrium, atmospheric pressure plasma discharges with bacteria has concentrated on the germicidal effects. Therefore, published results deal mainly with killing efficacy and little attention is given to physical mechanisms and biochemical pathways and their potential alterations when cells of microorganisms are exposed to the plasma. In this letter, an attempt to investigate the effects of plasma exposure on the biochemical pathways of bacteria is presented. In addition, using electron microscopy, we investigate if any gross morphological changes take place when cells are exposed to a lethal dose of plasma. We are testing the hypothesis that disruption of the cell membrane, sometimes to the point of cell lysis, is the mechanism whereby plasma kills cells. © 2002 American Institute of Physics. [DOI: 10.1063/1.1494863]

Recently, the ability to generate relatively large volumes of nonthermal gaseous discharges at atmospheric pressure allowed the emergence of exciting applications.1–4 Among these are the biological applications of atmospheric pressure glow discharges,4–7 which brought together plasma physicists and microbiologists. In addition to the basic scientific knowledge that such research is bound to contribute, plasmas are being considered as a potential means for the sterilization of biological and chemical warfare agents.4,5 The ability to generate nonthermal plasma discharges at pressures at or near 1 atm makes the decontamination process practical and inexpensive. In addition, the fact that the gas temperature in such discharges remains relatively low makes them use suitable for applications where medium preservation is desired. Unfortunately, the drive to develop a practical means of decontamination has led researchers to concentrate on the germicidal effects of plasmas with little attention given to their effects on fundamental physical and biological processes. Here we present an attempt to elucidate the effects of plasmas on the biochemical pathways of bacteria. In order to carry out such a study, cells of E. coli were exposed to a sublethal dose of cold plasma then seeded into various carbon substrates to evaluate any changes in substrate utilization relative to control cells. These changes are presumed indicative of plasma-induced changes in cell function. In addition, we used electron microscopy to investigate whether any gross morphological changes take place when cells of E. coli and Bacillus subtilis are exposed to a lethal dose of plasma. We are testing the hypothesis that disruption of the cell membrane is the mechanism whereby plasma kills cells.7,8

The plasma used in our experiments was produced by a resistive barrier discharge (RBD).9 Figure 1 shows the general configuration of the RBD. The plasma filled the gap (gap width = 4.5 cm) between two planar copper disk electrodes (diameter = 10 cm), one of which was covered by a high resistivity material. The electrodes were directly energized by a high voltage transformer delivering a voltage with a rms value up to 16 kV (at 60 Hz). The electrodes were contained within an enclosure in which a gas mixture of 97% helium and 3% oxygen was introduced. After the gas mixture was introduced, the plasma was ignited and maintained simply by adjusting the voltage source to the desired magnitude.

The biochemical impacts of a cold plasma generated by RBD on Escherichia coli were addressed with sole carbon substrate utilization (SCSU) experiments, using Biolog (Hayward, CA) GN2™ 96-well microtiter plates. The purpose of the SCSU experiments was to determine if exposure to plasma alters the heterotrophic pathways of the bacteria. It is presumed that any changes in metabolism would be indicative of plasma-induced changes in cell function. The Biolog GN2 plate is comprised of a control well and 95 other wells, each containing a different carbon substrate. Color development of a redox dye present in each well indicates utilization of that particular substrate by the inoculated bacteria.

FIG. 1. Configuration of the resistive barrier discharge.
The 95 substrates are dominated by amino acids, carbohydrates, and carboxylic acids. The extent and rate of color development for plasma treated cells and control cells were determined over five days at 24 hour intervals, by measuring the optical densities of each well with a microplate reader.

The optical densities (ODs) of the control wells were subtracted from the ODs of each of the 95 carbon substrates and an average well color development (AWCD) for each Biolog plate was calculated according to Garland and Mills.10 The AWCD was evaluated to determine whether an overall effect of the plasma treatment was observed. For the sublethal exposures administered in these experiments, no difference in AWCD between controls and plasma treated cells emerged. Next, the temporal pattern of color development for each carbon substrate was compared for the control cells and the plasma treated cells.

Although for most substrates no appreciable differences emerged, exposure to plasma was associated with significant changes in the utilization of some substrates. Figure 2 indicates an increased utilization (relative to the control cells) of several substrates (L-fucose, D-sorbitol, and D-galacturonic acid) by the plasma treated cells, and Fig. 3 indicates a decreased utilization of other substrates (methyl pyruvate, dextrin, and D, L-lactic acid). These changes were presumed indicative of plasma-induced alterations in some aspects of cellular functions (possibly enzyme activity).

To investigate any effects of cold plasmas on the morphology of bacteria we used scanning electron microscopy to inspect cells for gross physical damage following exposure to lethal doses. Representative Gram-negative (E. Coli) and Gram-positive bacteria (B. subtilis) were concentrated on filters, then exposed to plasma and subsequently examined microscopically. Figure 4 shows that cells of E. coli were heavily damaged following exposure to the plasma, to the point where membrane integrity was largely lost and their cytoplasm was distributed in clumps on the filter. Figure 5, on the other hand, shows that cells of B. subtilis exhibited no morphological change relative to the control cells.

Mendis, Rosenberg, and Azam8 theoretically predicted such an outcome on the basis of a disequilibrium between the forces exerted by the outward electrostatic tension and the tensile strength of the membrane. Mendis and co-workers8 claim that since the membrane of Gram-negative bacteria is irregular in structure, the outward elec-

![FIG. 2. Increased utilization of sole source carbon substrate by plasma treated E. coli cells. (a) L-fucose, (b) d-sorbitol, (c) D-galacturonic acid.](image1)

![FIG. 3. Decreased utilization of sole source carbon substrates by plasma treated E. coli cells. (a) Methyl pyruvate, (b) dextrin, (c) D,L-lactic acid.](image2)
trostatic tension (due to charge collection on the outer surface of the cells) is much higher than that in the case of Gram-positive bacteria, which exhibit a membrane having a smoother surface. This higher electrostatic tension can overcome the tensile strength of the membrane and induce its rupture.

This letter reports the effects of nonequilibrium atmospheric pressure plasma on heterotrophic pathways and cell morphology of bacteria. It shows that before cell destruction takes place, normal cell functioning can be altered by exposure to plasma. However, whether sublethal effects are reversible, cause irreversible damage (or mutation) to the cells, or ultimately lead to cellular death is unknown. In addition, the morphological effects of plasma exposure differed between the two species of bacteria tested. Membrane integrity of \textit{E. coli} was severely compromised following exposure to plasma, but no apparent morphological effects were manifested on \textit{B. subtilis}.

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