Urinary Tract Infection Risk Assessment By Non-Thermal Plasma In Iraqis Patients

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**ABSTRACT**

**Background:** In the present study used device jet plasma needle with atmospheric pressure which generates non-thermal plasma jet to measure treatment potent with plasma against pathogenic bacteria founded in UTI was inactivated with plasma at 10 sec.

**Objective:** This work included the application of the plasma produced from the system in the field of bacterial sterilization, where sample of Gram-negative bacteria (Escherichia coli) were exposed to intervals (1-10) second.

**Methods:** The work were used in this study obtained from studying 100 urine samples, the age of patients ranged between 10 years to 60 years. They were 60 females and 40 males. These samples were cultured on culture media to isolate bacterial colonies. After that, bacteria were identified by means of highly specific investigations. Escherichia coli, plasma needle treatment is applied on bacteria through sterilization, and adhesion.

**Results:** It was found that the percentage of the killing of Gram-negative bacteria (E.coli) was 100% at (10) second, also decreasing bacterial adhesion on epithelial cells, where numbers adhesion bacterial with uroepithelial cells decrease after treatment with plasma needle.

**Conclusion:** From this work, it has been observed that applied voltage, distance between plasma needle and treatment model as well as time treatment effect on inactivation bacteria and sterilization, also it effect on decreasing bacterial adhesion on epithelial cells, where numbers adhesion bacterial with uroepithelial cells decrease after treatment with plasma needle.

**Keywords:** Urinary Tract Infection, plasma needle, adhesion, E. coli bacteria

Al-Kindy College Medical Journal 2017: Vol. 13 No. 1

Page: 86-91

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Received 15th Feb 2017, accepted in final 30th March 2017

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killing bacteria. This makes these plasmas very useful for various biological and medical applications, such as: sterilization of medical instruments, decontamination in biological warfare and air filters in hospitals [8]. The plasma needle is a type of non-thermal atmospheric glow discharge, it has a single electrode configuration and is operate by different noble gas (He-Ar), important properties of this type of plasma are that it operate at near room temperature, the plasma does not cause any thermal damage to articles it comes in contact with. This characteristic was open up the possibility to use this plasma for treatment of the heat sensitive materials. Atmospheric pressure discharge plasma is of great interest because of their low costs and simplified operation [10]

Non-equilibrium plasma at atmospheric pressure finds numerous biological and bio-medical applications thanks to their reactive nature. It has been tested on large variety of bacteria, spores, viruses for their sterilization and interactions of plasma with live tissue e.g., skin disinfection, blood coagulation, wound healing, density [11,12]. In bio-decontamination by plasma, it is crucial to understand the role of various mechanisms involved. The significant mechanisms depend on the plasma composition (gas), temperature, treated microorganisms and the environment (air, water, surfaces, etc.) [13].

The most common cause of infection is Escherichia coli, though other bacteria or fungi may rarely be the cause [15]. Kidney infection, if it occurs, usually follows a bladder infection but may also result from a blood-borne infection [17].

In women, they are the most common form of bacterial infection [16]. Up to 10% of women have a urinary tract infection in a given year and half of women having at least one infection at some point in their lives [17]. The bacteria that cause urinary tract infections typically enter the bladder via the urethra. It is believed that the bacteria are usually transmitted to the urethra from the bowel, with females at greater risk due to their anatomy. After gaining entry to the bladder, E. CoE are able to attach to the bladder wall and form a biofilm that resists the body's immune response [17].

Rates of asymptomatic bacteria in the urine increase with age from two to seven percent in women of child bearing age to as high as 50% in elderly women in care homes [15]. Rates of asymptomatic bacteria in the urine among men over 75 are between 7-10%. Asymptomatic bacteria in the urine occurs in 2% to 10% of pregnancies [13,17].

Effect adhesions on Epithelial cells: Adhesions are cell-surface components or appendages of bacteria that facilitate bacterial adhesion or adherence to other cells or to inanimate surfaces. Adhesions are a type of factor. Adherence is an essential step in bacterial pathogenesis or infection, required for colonizing a new host [18]. Most fimbriae of gram-negative bacteria function as adhesions, but in many cases it is a minor subunit protein at the tip of the fimbriae that is the actual adhesion. In gram-positive bacteria, a protein or polysaccharide surface layer serves as the specific adhesion. To effectively achieve adherence to host surfaces, many bacteria produce multiple adherence factors called adhesions [19].

Epithelial cells cover the whole surface of the body. It is made up of cells closely packed and ranged in one or more layers. This tissue is specialized to form the covering or lining of all internal and external body surfaces. An epithelial cell that occurs on surfaces on the interior of the body is known as endothelium. Epithelial cells are packed tightly together, with almost no intercellular spaces and only a small amount of intercellular substance. Epithelial cells, regardless of the type, is usually separated from the underlying tissue by a thin sheet of connective tissue; basement membrane. The basement membrane provides structural support for the epithelium and also binds it to neighbouring structures [20].

Urinary tract infections UTI, are the most common disease in human, adhesion process is first step to establish the infection, where bacteria is double after adhesion to arrive of forming colonization, for bacteria appliance process of adhesion and isolated bacterial power in adhesion depends on saturate extent and receivers concentration within epithelial cells, same method, bacteria which contains adhesion processes such as Fimbria ability on adhesion with bladder cells consequently cystitis, while loses bacteria of structures can not adhesion consequently excretion with urine. Bacterial adhesins provide species and tissue tropism. Adhesins are expressed by both pathogenic bacteria and saprophytic bacteria. This prevalence marks them as key microbial virulence factors in addition to a bacterium's ability to produce toxins and resist the immune defenses of the host. Mature Fim H is displayed on the bacterial surface as a component of the type 1 fimbrial organelle [19].

The majority of bacterial pathogens exploit specific adhesion to host cells as their main virulence factor. A large number of bacterial adhesins with individual receptor specificities have been identified. Many bacterial pathogens are able to express an array of different adhesins. Expression of these adhesins at different phases during infection play the most important role in adhesion based virulence.

Methods

Experiment setup

Plasma needle designed with diameter 1mm from interior, this needle constitutes cylindrical tube made from glass material with length 100mm interior this glass tube, put other cylindrical tube made from iron material with external diameter 2.7 mm, this tube connect to anode from high voltage power supply about 9.6kV peak to peak, applied power was lasting of electrical discharge which calculated from simultaneous values of voltage and current about 15 watt and applied frequency 33khz , it through pass argon gas where discharge between electrode and space through needle hole where plasma generation outside from hole, figure (1) shows plasma needle.

![Figure (1): plasma needle](image_url)

Methods: Hundred patients (100) complaining from different clinical signs and symptoms were admitted to Ramadi Teaching Hospital for seeking medical advise were included in the present study. Midstream urine samples (MSU) were obtained from patient with urinary
Urinary Tract Infection

Mohammed Ubaid Hussein and Rana Talb Mohsen

College Medical Journal 2017: Vol.13 No. 1
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Urinary tract infections (UTIs), were examined bacteriological. These samples were collected from patients visiting the urology department -Ramadi Teaching Hospital. The results of this work were obtained from studying 100 urine samples. These samples were taken from patients complaining from signs and symptoms of urinary tract infections (UTIs) they were 60 females and 40 males. The technical detail were described in a methodological study, bacteria were transferred from deep agar subculture where they had kept after isolation to MacConkey agar plates. After growth for 16-48 hrs at 37°C in Brain Heart Infusion (BHI) broth, the bacteria were centrifuged and resuspended in Phosphate Buffer Saline (PBS) pH (7.4). Uroepithelial cells (UECs) were obtained from the sediment of the midstream morning specimen of healthy females without bacteriuria. They were washed and resuspended in PBS [21,22,23]. A drop of the final UECs suspension was deposited on a glass slide, air-dried, heat fixed and Gram stained. A control slide with no attached bacteria was stained at the same time. Examination of the slide was done by light microscopy under oil immersion to demonstrate the attachment bacteria to UECs [24,25]. Two tubes, one before plasma needle treatment, other after plasma needle treatment. The adhesion for each bacterial isolate was estimated as the mean number of bacteria adhering to many epithelial cells. The results were compared with the results of control group.

Also, bacterial samples were grown in brain heart infusion broth at 37°C for 20 hour. From this suspension, suspension with a number of bacteria about 10^8 (CFU/ml) were made as determined by 0.5 McFarland standard and spectrophotometric assays. The resuspended culture was serially diluted to 1:10, 1:100, 1:1000, 1:10000 of the original. The last diluted suspension was used in this experiment to evaluate the effect of plasma needle system on bacteria. To prevent microbial contamination during experiment, plasma system was placed in a sterile condition. In this experiment 20µL drop of prepared bacterial suspension were placed on agar surface. This volume was selected as it spread to ≈ 1 cm^2 over the agar surface; thus, the area covered by the bacterial sample drop was entirely within the area covered by the insulated plasma needle.

Results: The plasma needle effects, on bacteriology through samples taken of patients, methods sterilization and adhesion were shown. The results of this work were obtained from studying 100 urine samples, the age of patients ranged between 10 years to 60 years. They were 60 females and 40 males, as in figure (2).

Figure (2): Sex distribution (female & male) for patterns those help in these experiments.

The glass petri dishes were placed on the ground one by one and exposed to plasma needle for subsequent times (1-20) s. In addition to exposure samples, control samples were inoculated but not exposed to plasma needle. After the treatment, the drop was spread over the entire agar surface and incubated at 37°C for 24 h. After incubation, the colony forming units (CFU) were counted in order to check the efficiency of bacterial inactivation using plasma needle system.

1. Treatment of Bacterial Sample: *Escherichia coli* were selected to evaluate the inhibitory effect of laboratory-made non-thermal atmospheric pressure plasma (plasma needle) on their growth using brain heart infusion agar culture media, the results shown in figure (3) for *Escherichia coli* Bacteria. The figures shows the reduction in colony forming unit (CFU) as increasing the exposure time as shown figure (4).

Figure (3): *E.coli* growth samples after exposure to plasma needle system
A: control, B: 2 second, C: 8 second, D: 10 second

Figure (4): Relation ship between number of survivals (CFU) & exposure time (second) of *E.Coli* bacteria
The germicidal effect of plasma system on *E.coli* isolates were analyzed using a germicidal efficiency equation as follows:

\[
\text{Germicidal efficiency} = \left(\frac{N_0 - N_t}{N_0}\right) \times 100\%
\]

- **N** = CFU of non-treated bacteria (control).
- **N**<sub>t</sub> = CFU of treated bacteria.

Results show that plasma needle treatment during 10 seconds effectively sterilized *E.coli* with a 100% percentage, as shown figure (5).

**Figure (5): Germicidal efficiency of plasma needle on *E.coli* at different plasma treatment time.**

2. **Adhesion**

This study shows effect the plasma needle on pathogenic bacteria causing UTI, and role it in adhesion process of this bacteria on epithelial cells of UTI. One type of bacteria was included in this study *E.Coli* bacteria after isolated from clinical samples of patient suffering UTI.

*E.Coli* bacteria polysaccharides play important role in capability these bacteria on adhesion with epithelial cells, for wallet important affect through first steps of disease these bacteria, where gets contact them and between epithelial cells producing epithelial material, also affect the presence of the wallet in process of building proteins necessary of structure bacterial adhesion.

The adhesion for each bacterial isolate was estimated as the mean number of bacteria adhering to many epithelial cells. *E.Coli* bacteria was selected to evaluate the inhibitory effect of laboratory-made non-thermal atmospheric pressure plasma (plasma needle) on their growth. The results were compared with the results of control group before treatment by plasma needle.

Figure (6) shows the mean values of No. of adherent *E.Coli* bacteria to epithelial cells before & after treatment with plasma needle.

**Figure (6): Mean values of No. of adherent *E.Coli* bacteria to epithelial cells before & after treatment with plasma needle.**

Figure (7) shows the mean values of No. of bacteria attached to epithelial cells before & after treatment with plasma, where it decreases after treatment at applied voltage 10.6 kV, distances between plasma needle and substance (treatment model) were 2 and 1 cm and frequency 33 kHz through period 1.30 min.

**Figure (7): Mean values of No. of adherent *E.Coli* bacteria to epithelial cells before & after treatment with plasma needle.**

Increasing applied voltage, treatment time and decreasing distances between plasma needle and substance, effect on decreasing No. of bacteria attached to epithelial cells on *E.Coli* bacteria.
**Discussion:** Bacterial adhesion process causes of UTIs through two stages, first recognition phase initial, this stage depends on pili especially Type 1-pili and Type p-Fimbria with interaction it with epithelial cells and covers it from receptors. Second stage, adhesion more resting where other surface structures involve with it such as O-antigen or lipopolysaccharide founded with outer membrane of Gram negative bacteria where bacteria has this structure led to increasing ability it on adhesion.

Most gram negative bacteria isolated which causes cystitis, fimbria (Type1 Fimbria) ability to connect by mannose units in mucous cells or the cell membranes of host.[19,27]

Plasma needle effect in blocking mannose units existing with epithelial cells, thus reduces the bacterial adhesion rate with epithelial cells. As for *E.Coli bacteria*, cell wall contains Teichoic acid helps adhesion[10], where role of plasma on decreasing adhesion with epithelial cells isolated from UTI, consequently these results show that plasma refuses bacteria from adhesion with bladder wall, consequently inactivation process adhesion of bacteria with epithelial cells of UTI. The specific mechanism for the plasma effect on epithelial cells is similarly unclear. Cold plasma produces long living (O$_3$,NO,H$_2$O$_2$) and short lived (OH, O electronically excited) neutral particles and charged particles(ions and electrons). All of these could be toxic to cells, induce low levels of cell membrane damage and potentially change intercellular signaling pathways. Specific plasmas can be created to produce either neutrals or charged particles in order to elucidate pathways. Specific plasmas can be created to produce either neutrals or charged particles in order to elucidate the critical mechanism. Charged particles can play a very significant role in the rupture of the outer membrane of bacterial cells.

**Conclusions:** There was laboratory plasma needle system showed germicidal effect on Gram-negative bacteria (*E.Coli*), charged particles found in plasma can play a very significant role in the rupture of the outer membrane of bacterial cells, also change of applied voltage and distance by sterilization method, as shown. Plasma needle treatment is applied on bacteria through sterilization, and adhesion. From this work it has been observed that applied voltage, distance between plasma needle and treatment model as well as time treatment effect on inactivation bacteria and sterilization so it effect on decreasing bacterial adhesion on epithelial cells, where numbers adhesion bacterial with uroepithelial cells decrease after treatment with plasma needle.

**Acknowledgments:** I would like to express special words of thanks with deepest appreciation of college medicine and Clinical Laboratories in Al-Ramadi General Hospital, also the Staff working in these Laboratories.

**References**


