Effect of Silver Nanoparticles on Common Bacteria in Hospital Surfaces

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ABSTRACT

Background: One of the most important causes of complications and mortality in medical centers are nosocomial infections. Disinfection of hospital surfaces is essential element for ensuring that infectious agents are not transmitted to patients. Alcohol-based and chlorine-based disinfectants have unfavorable properties. Given that the antimicrobial effect of heavy metals such as silver is recognized as a viable option for eliminating bacteria, the exploration of nanotechnology in this context has been described in this study. Nanotechnology uses both science and technology to produce new materials with nano-scales.

Objectives: The effect of silver nanoparticles on some common hospital bacteria has been studied in this research.

Patients and Methods: We have selected nine patients’ metal file covers and following sterilizing, we have infected them with one of these bacteria; Staphylococcus aureus, Pseudomonas aeruginosa and Bacillus cereus. Then, the infected surfaces have been disinfected with different dilutions of silver nanoparticles. Sampling and culturing has done following four specific intervals. Afterwards, the colonies that developed have been counted and compared.

Results: All of the three dilutions of silver nanoparticles could bring the colony count out of 7.5x10^6 to less than 100 which indicate more than a 99 percent reduction. No remarkable difference of the three dilutions of disinfectants was observed in reducing the colony count in 5, 15, 30 and 60 minute disinfection intervals (P value > 0.05). The effect of constant dilution of silver nanoparticles on the reduction of organisms varied in response to the time of disinfection (P value < 0.05).

Conclusion: Silver nanoparticles had appropriate effects in all three types of dilutions and allowing for a more protracted contact time has given significantly better results.

Keywords: Disinfection; Nosocomial Infection; Disinfectant.
1. Background

Nosocomial infections are one of the most important causes of complications and mortality in medical centers. These infections have been considered as the sixth leading cause of death in US and Europe. As well, they contribute 10-50 billion dollars to the economic burden annually (1). On the other hand, devices used for patient care were divided in to three groups based ontheriskof being infectious: critical items, semi-critical items and non-critical items.

Critical items enter sterile tissue and the vascular system. Therefore, if they are contaminated with microorganisms such as bacterial spores there is a high risk of infection. Therefore these items should be sterile (2).

Semi-critical items have contact with mucous membranes and skin that has been compromised. So, there is a necessity to disinfect microorganisms with a high-level chemical disinfectant (2, 3). Non-critical items have contacts with skin, but not with mucous membranes. They are divided in to two groups: non-critical patient care items and non-critical environmental surfaces (2, 3).

Examples for non-critical patient care items are the bedpan, blood pressure cuff and computer system. In fact, they are not dangerous and do not infect patients with infectious agents (4). Non-critical surfaces include: the patients’ bed rail, the table beside patients’ bed, that have repeated contact with hands and can potentially contribute to transmitting secondary infectious agents by infecting hospital staffs’ hands and other medical items which have contact with the patients (5, 6).

Disinfection is a process that eradicates some or the entire pathogen microorganisms except bacterial spores on inanimate surfaces (7). Disinfectants threaten healthcare workers’ health from mucosal irritation to death (8). However, toxicity degrees are different (9, 10) but all disinfectants must be used with precautions (11). Based on the efficiency grade, disinfectants are divided in to three levels: high, medium and low (12).

Comprehensive studies on anti-septic (13) and disinfectants (14) clearly indicate the anti-microbial effect of heavy metals (15, 16). Silver was used for prevention of newborns’ conjunctivitis, as a topical treatment for burn trauma and impregnating a different catheter (17).

Nano-technology is a new developing context using science and technology with the aim of producing new materials with nano-scales (18). The term nano-technology was first created by Professor Nario Taniguchi in Japan in 1934 that described producing precise materials in nano-scales (19). Different types of nano materials include titanium (20), gold (21), alginate (22) and silver have been studied however the antimicrobial effect of silver nanoparticles demonstrated that it is the most effective substance against bacteria, viruses and eukaryotic microorganisms (23). This superior result is because of the very large surface that causes better contact with microorganisms. Nanoparticles adhere to the cell membrane and permeate in to the bacterium (24-26).

Using silver nanoparticles as a disinfectant may provide concerns about the risk of silver toxication (Argyria) and its toxicity for the body cells (27). Research in this field indicate that silver ions or nano-particles can be applied for the treatment of burns, dental materials, sunscreen lotions and so on. It has low toxicity for human cells (24).

2. Objectives

Alcohol-based and chlorine-based disinfectants have adverse side effects such as irritations, skin allergies and inhalation risks as well as an annoying and offensive odor. The disinfection effect of these substances on the multitude of existing resistant microorganisms is controversial. For this reason, we studied the effect of the different dilutions of silver nanoparticles on some of the common hospital organisms at specific intervals. Provided that the effect of silver nanoparticles proves to be an effective, viable alternative to traditional hospital disinfectants, we recommend that this substance be used in their stead.

3. Patients and Methods

In a single blind interventional study, we have selected nine patients’ metal file covers as representative of common hospital surfaces. We sterilized these surfaces with an autoclave before beginning the intervention. Subsequently, we collected swab samples from those surfaces and cultured them to ensure that we have reached the objective of sterile surfaces. If there is any culture growth on these surfaces, that surface will be excluded from the study and the test as described will be repeated.

3.1. Contaminating Surfaces

We have dedicated four equal 2x2cm squares on every sterile metal cover (a, b, c, d). Sterile surfaces were infected with 50microliters of the selected organisms under study with concentration of 0.5 McFarland. Regarding that 1 milliliter of 0.5 McFarland contains 1.5 x 10^8 bacteria, we have used 7.5 x 10^4 bacteria to contaminate every 2x2cm surface. Bacteria were then extracted from the positive clinical samples of the hospital laboratory and their strains have been verified by standard biochemical tests. We have dedicated 3 file cover surfaces for each kind of bacteria with the aforementioned four 2x2cm squares on each surface.
The culture media contained the following kinds of bacteria colonies: methicillin resistant *Staphylococcus aureus* MRSA (as a common nosocomial infection and colonization Gram-positive bacteria), *Pseudomonas aeruginosa* (with antibiogram resistant to ceftazidime and ceftizime as common hospital Gram-negative bacteria) and *Bacillus cereus* (as common nosocomial infection and colonization sporulative bacteria). A 0.5 McFarland suspension of each culture media was prepared and every four squares (a, b, c, d) of file covers have been contaminated with the amount of bacteria (50 microliter) as follows: files 1-2-3 (s1, s2, s3) contaminated with *S. aureus*, files 4-5-6 (p1, p2, p3) with *P. aeruginosa* and files 7-8-9 (b1, b2, b3) with *B. cereus*.

### 3.2. Disinfection of Contaminated Surfaces

All squares on the nine metal covers have been disinfected with swab which was saturated with one of the three silver nanoparticles’ dilutions (100ppm, 200ppm, 300ppm). To break it down, we had four squares on s1, p1, b1 with 100ppm silver nanoparticles, four squares on s2, p2, b2 with 200ppm silver nanoparticles and four squares on s3, p3, b3 with 300ppm silver nanoparticles. It was assumed that PPM means part per million, and silver nano-particles were diluted with sterile water to these dilutions.

### 3.3. Sampling and Culturing

Following 5 minutes, the square (a) of each metal cover had been sampled by a sterile swab and this swab was put in a test tube containing normal saline. Until then, it had been transferred to the culture media. Sampling was repeated for b, c and d squares after 15, 30 and 60 minutes, respectively. The encoded test tubes had been put at a microbiologist’s disposal. The microbiologist was unaware of the bacteria type, the disinfectant substance used in each test tube and the duration time was not indicated. It had been transferred to the culture media. All squares on the nine metal covers have been disinfected with swab which was saturated with one of the three silver nanoparticles’ dilutions (100ppm, 200ppm, 300ppm). To break it down, we had four squares on s1, p1, b1 with 100ppm silver nanoparticles, four squares on s2, p2, b2 with 200ppm silver nanoparticles and four squares on s3, p3, b3 with 300ppm silver nanoparticles. It was assumed that PPM means part per million, and silver nano-particles were diluted with sterile water to these dilutions.

### 3.4. Results

No microorganisms grew in the first samples collected from sterile surfaces. Therefore no surface excluded from the study. We calculated the average colony count divided by the type of microorganism and the total average of counted bacteria for each specific time period divided by the disinfectant dilution. The results can be seen as recorded in Table 1.

All the three dilutions of silver nanoparticles had definite positive effects on the bacteria under study as can be seen in Table 1. The silver nanoparticles could bring the colony count out of 7.5x10⁶ to less than 100 which indicates more than 99 percent reduction. No remarkable difference was indicated in comparing different dilutions of silver nanoparticles on the total of all microorganisms which were applied for contaminating surfaces in a specific time (P value > 0.05 in all four specific times, Figure 1).

### Table 1.

Mean Number of Calculated Studying Bacteria Colony Count in the Culture Media Following Times of Disinfection

<table>
<thead>
<tr>
<th>Disinfectant</th>
<th>Silver Nanoparticles 100 ppm</th>
<th>Silver Nanoparticles 200 ppm</th>
<th>Silver Nanoparticles 300 ppm</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bacteria/</td>
<td>Time</td>
<td>5</td>
<td>15</td>
</tr>
<tr>
<td>MRSA³</td>
<td>s1</td>
<td>10/33</td>
<td>0/33</td>
</tr>
<tr>
<td></td>
<td>s2</td>
<td>34</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>s3</td>
<td>46/66</td>
<td>19/33</td>
</tr>
<tr>
<td>Mean of T.⁴</td>
<td>62/38</td>
<td>6/77</td>
<td>0/77</td>
</tr>
</tbody>
</table>

³Abbreviations: Mean of T, Mean of Total of all Bacteria; MRSA, Methicillin Resistant *Staphylococcus*; PA, *P. aeruginosa*; BC, *Bacillus cereus*
Although 300ppm silver nanoparticles showed a significant efficacy in 15, 30 and 60 minutes, the difference in efficacy between intervals had no statistical meaning, as mentioned beforehand. By allowing longer contact time with each dilution of disinfection in our study, we had a significant colony count reduction compared to the former specific contact time (P value < 0.05). 100ppm, 200ppm and 300ppm silver nanoparticles had respectively P value of 0.01, 0.03, and 0.01. The calculated average total number of colony count for each three types of bacteria was calculated comparatively with definite disinfection time in culture media divided by the dilution of the disinfectants as is indicated in the Figures 2, 3, 4.

5. Discussion

Following five minutes all disinfectants reduced the S. aureus colony count significantly (more than 99%), but none of them could bring the S. aureus colony count to zero. This rate was achieved following 30 minutes of disinfection with 300ppm silver nanoparticles.

For two other dilutions of silver nanoparticles, this rate was possible after 60 minutes of disinfection. Thus 300ppm effect of silver nanoparticles on reducing S. aureus colony count is faster than the two other ones, but no remarkable difference of the three dilutions of disinfectants was observed in reducing S. aureus colony count in 5, 15, 30 and 60 minute intervals of disinfection (P value >0.05).

This proper effect of silver nanoparticles on S. aureus has studied in some other research. A 100% reduction of S. aureus was indicated following 48 hours contact time of silver nanoparticle-coated masks in Li study (16). Plastic catheters coated with silver nanoparticles were contaminated with S. aureus and 100% effect of eradicating was observed following 72 hours in David Roe and colleagues study (28). 12.5ppm silver nanoparticles in a culture media showed a 100% reduction of S. aureus colony.
count following 5 minutes in Ghazvini and colleagues’ study, but the lower dilutions were not efficient. The effective dilution of Ghazvini’s study was lower than in the present study. This difference can be explained by the fact that Ghazvini has set silver nanoparticles directly in contact with *S. aureus* in a culture media (29).

Antimicrobial effect of silver nanoparticles on the growth of staphylococcus aureus was reported to be very insignificant in Kim’s study. Kim assessed the antimicrobial effect of silver nanoparticles in a culture media, not as a disinfectant of surfaces. They studied low dilutions of this substance with the unit of nanomole. MIC more than 33nM of silver nanoparticles for *S. aureus* and MiC more than 6.6nM for Escherichia coli were measured (Every 1000 nanomole equal to 1ppm). These items may justify the differences between Kim’s study and ours (30). We had a zero *P. aeruginosa* colony count following 5 minutes of disinfection with silver nanoparticles and more than 99% reduction after 5 minutes. It means that three dilutions of silver nanoparticles have similar effects on *P. aeruginosa* colony counts of hospital surfaces after a specific disinfection time (P value > 0.05).

There are other studies about the disinfection efficacy of silver nanoparticles on *P. aeruginosa* which had various results. Our study has similar findings to Lara and colleagues’ research in Mexico which was about bactericidal efficacy of silver nanoparticles on *P. aeruginosa* (31). 100ppm silver nanoparticles indicated a 100% reduction of *P. aeruginosa* colony count following 15 minutes in culture media in Ghazvini’s study which was quite similar to ours (29).

David Roe and colleagues’ study represented 70% efficacy after 72 hours contact time. They had experimented on plastic catheters coated with silver nanoparticles contaminated with *P. aeruginosa*. Their result was not similar to ours; as well, it was unlike other mentioned studies. Considering the different method of this study from other mentioned studies, this difference could be explainable (28). *B. cereus* growth reduction was 100% after 60 minutes disinfection with 300ppm silver nanoparticles as is observed in the results. Actually, growth reduction of other dilutions (100ppm and 200ppm) was rather high (more than 99%) in 60 minutes. However none of them could bring *B. cereus* colony count to zero. Although the 300ppm silver nanoparticles’ efficacy was much more than the others, this difference was not significant (P value > 0.05).

Ghazvini’s research also studied the effectiveness of silver nanoparticles on *B. cereus*. The maximum dilution applied in that study was 200ppm which could not bring *B. cereus* colony count to zero in five, 15, 30 minutes and 24 hours. Although this dilution could indicate more than 5log reduction which was 99% of the mentioned bacteria colony count. Considering the fact that we gained almost similar results of this dilution in our study and 300ppm could reduce 100% of colony count following 60 minutes seems to indicate that both studies demonstrated similar efficacy of silver nanoparticles (29).

As revealed, there was a good effect with no remarkable difference in comparing different dilutions of silver nanoparticles on all microorganisms which were applied for contaminated surfaces in a specific time (Figure 1). This proper effect of silver nanoparticles could be highly suitable to decrease all kinds of organisms, considering the fact that hospital surfaces are often infected with multiple organisms. 300ppm silver nanoparticles could bring the number of all bacteria to zero after 60 minutes disinfection. This number was near zero after 30 minutes. Regarding the fact that we had sporulative Bacillus in this study, it would seem that 300ppm silver nanoparticles could be recommended as a high level disinfectant.

Statistical analysis indicated the efficacy of silver nanoparticles disinfection on decreasing colony count of *S. aureus* and *P. aeruginosa* has a remarkable difference with decreasing of *B. cereus* colony count. This difference is observed in all studied dilutions but statistical comparison was not possible, because of having low number of samples and zero number of colony counts in at specific time intervals. Considering the fact that bacillus cereus is sporulative probably the efficacy of different dilutions of silver nanoparticles on non-sporulated Gram-positive and Gram-negative bacteria is greater than sporulated ones, such as bacillus cereus.

In David Roe and colleagues’ study on the plastic catheters impregnated with silver nanoparticles, the efficacy of this disinfectant on *S. aureus* after 72 hours contact time was 100% which was more than 70% efficacy on *P. aeruginosa*. There were no statistical comparisons (28). Our research results have many similarities to Ghazvini and colleagues’ study in Mashad, Iran. We also found the strong bactericidal effect of silver nanoparticles against Gram-positive bacteria like *S. aureus* and Gram-negative bacteria like *P. aeruginosa* and lower efficacy of this substance on sporulative bacteria like *B. cereus* compared to the effects on *S. aureus* and *P. aeruginosa* (29).

The effect of constant dilution of silver nanoparticles on the reduction of hospital surfaces’ organisms (*S. aureus, P. aeruginosa* and *B. cereus*) is different at various times of disinfection. This positive result of extended time on silver nanoparticles’ disinfecting *S. aureus* and *P. aeruginosa* revealed in Ghazvini’s study. But, this issue was not verifiable for *B. cereus*. As was mentioned before, Ghazvini’s research did not study surfaces, so there is a definitive difference between that study and ours (29).

The strong efficacy of silver nanoparticles on reducing bacterial organisms’ growth (more than 99%) is the result of this study which mirror similar achievements of prior studies which support the significant viability of silver nanoparticles as a viable alternative for hospital common area disinfectants. Silver nanoparticles of lower dilutions
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(100pm) can also be applied upon susceptible surfaces without any worry of having less effectiveness as a disinfectant. Therefore, there appears to be no need to use a higher concentration of silver nanoparticles which makes them a more affordable alternative.

Acknowledgements

The authors would like to thank the personnel of infectious diseases ward of Ahvaz Razi Hospital and laboratory of microbiology of Jundishapur University of Medical Sciences for their help and also appreciate the Nano Nasb Pars Company that had provided us silver nanoparticles free of charge for the study.

Financial Disclosure

None Declared.

Funding Support

This study was supported by research center of infectious and tropical diseases of Ahvaz Jundishapur University of Medical Sciences, Iran.

Authors' Contribution

None declared.

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