Comparative evaluation of anti-bacterial efficacy of silver nanoparticle, nanoparticle calcium hydroxide and calcium hydroxide against enterococcus faecalis biofilm- An In vitro study

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Abstract

The purpose of this study was to evaluate the antibacterial efficacy of silver Nanoparticles, Nanoparticle calcium hydroxide and calcium hydroxide as intracanal medicament against Enterococcus faecalis bio film formed on root dentin. The Minimal inhibitory concentration test and Anti microbial test were conducted for silver nanoparticle, Nanoparticle calcium hydroxide. Teeth were inoculated with E. faecalis strains for 1 month to establish a standard mono species bio film model. Biofilm formation is confirmed using SEM. The intracanal medicaments were placed inside the canal and colony forming units (CFU) were counted before and 7 days later the placement of the medicament. MIC for AGNP is 25µg/200µl and for NCH is 12.5µg/200µl. The Antimicrobial test conducted shows zone of inhibition about 11mm and 14mm for AGNP and NCH respectively at 500µg. The calcium hydroxides do not show any antibacterial activity at this concentration. In antibacterial efficacy test using CFU count, the AGNP and NCH shows 75% and 61% of antibacterial efficacy respectively whereas calcium hydroxide has exhibited just 11% of antibacterial efficacy. Silver Nanoparticles and nanoparticle calcium hydroxide had statistically significant difference in the reduction in the number of colonies when compared to calcium hydroxide by post hoc turkey test and p-value of <0.01 was considered as significant in this study.

Keywords: Silver Nanoparticles; Calcium Hydroxide; Intracranial Medicaments; E. Faecalis

1. Introduction

The basis of endodontic treatment depends on identifying and eliminating the causative factors in the development of apical periodontitis so that optimal healing can be achieved. The role of bacteria and their byproducts in the pathogenesis of apical periodontitis has been clearly established. An improved prognosis has been shown for the teeth after a negative culture has been obtained compared with teeth having a positive culture at the time of obturation. Therefore, the prime objective of treatment is to eliminate bacteria and sources of nutrient supply from root canal system. [1] It was well known that after the Instrumentation of the infected pulp space, some type of further disinfection would be required to enhance the chances for treatment success. The proven step to take is to apply an effective antimicrobial agent in the root canal for a predetermined time period to further eradicate the remaining bacteria. Enterococcus faecalis is the predominant micro-organism and occasionally the only species detected in root canals of teeth associated with persistent periapical lesions. It is hardy microbe that possesses certain virulence factors including lytic enzymes, cytolsin, aggregation substances, pheromones, and lipoteichoic acid. E. faecalis is able to invade dental tubules and remain viable within the tubules for prolonged period of time, adhere and form biofilm on dentin under different environmental conditions, resist intracanal disinfectants, survive harsh conditions within root filled teeth. Historically, efforts to eliminate E. faecalis and its concomitant biofilm have been somewhat limited while using commonly used root canal disinfectants. [2].

Since its Introduction as an intracanal medicament in 1920 (HERMANN 1920) calcium hydroxide has been widely used in Endodontics. It is a strong alkaline substance, which has a Ph of approximately 12.5. Various biological properties have been attributed to this substance, such as Antimicrobial activity (Bystrom et al 1985), tissue dissolving ability (Hasselgren et al 1988, Andersen et al 1992), because of such effects calcium hydroxide is the most commonly used Intracanal medication during root canal procedures. Its antibacterial property is generally related to the release of hydroxyl ions, which produces the lethal effects on bacterial cells including protein denaturation and damage to the bacterial cytoplasmic membranes and DNA. However, calcium hydroxide can be inactivated by dentin exudates from periapical area and microbial biomass. Recently, much attention has been focused on Nanoparticle in many health care fields. With regard to their high surface-to-volume ratio, Nanoparticles are one of the most effective antibacterial agents. Silver nanoparticle (AgNPs) has been applied in many health care fields because of their broad spectrum bactericidal and virucidal properties. AgNPs have high surface- area-to-volume ratio.
and unique chemical and physical properties which results in increased reactivity. AgNPs shows multiple antibacterial mechanisms such as adherence and penetration into bacterial cell membrane and cell wall permeability. AGNPs with size in the range of 10-100 nm showed powerful bactericidal potential against both gram positive and gram negative bacteria. [4] Similarly calcium hydroxide is a commonly used intra canal medicament but it lacks in ability of completely eliminating Enterococcus faecalis. Nanoparticle of calcium hydroxide (NCH) would penetrate dentinal tubules deeper and at a greater concentration than calcium hydroxide. NCH have superior anti microbial activity against E. faecalis compared to conventional calcium hydroxide in culture media as well as dentin tubules. So to evaluate the antibacterial efficacy of Nanoparticles, this study uses silver nanoparticle and Nanoparticle calcium hydroxide along with calcium hydroxide against mature E. faecalis biofilm.

2. Materials and methods

2.1. Standardization of the specimens

32 single rooted teeth (maxillary and mandibular anterior and mandibular premolars) were selected and stored in 0.5 % Thy- mol. Teeth are sectioned 3mm above CEJ and prepared with Protaper up to F3 maintaining patency. Smear layer removed (2 ml – 6% NAOCL, 2 ml -17% EDTA, 2ml – 6% NAOCL with positive pressure irrigation using side vented needles). Teeth are wrapped in moist cloth and sterilized in autoclave. Teeth transferred to sterile vials containing 20ml sterile BHI (brain heart infusion) and kept in incubator at 37 o C for 48 hrs to check efficacy of sterilization procedures.

2.2. Cultivation of enterococcus faecalis and specimen inoculation

E.faecalis colonies are grown on BHI agar plates were suspend- ed in 20 ml sterile BHI broth for 8 hr. Five drops of BHI broth is added to 20 ml broth for 4 hours. Bacterial suspension are adjusted to 1.5 X 10 ⁶ CFU / ml turbidity. 20 micro liters of bacte- rial inoculums added to vials containing the teeth suspended in sterile medium. Tubes closed and stored on 37 o C for 30 days to ensure adequate bacterial penetration in to dentinal tubules. Half of the inoculums broth for all specimens was replaced with 20 ml of sterile BHI medium every 5 days. Biofilm formation is confirmed using SEM.

2.3. Samples and microbiological analysis

2.3.1. Methodology for antibacterial effectiveness of intracanal medicament mixtures against E. faecalis

silver nanoparticle, nanoparticle calcium hydroxide at different concentration of 500,250,125,50,12.5 µg/ 200µl is taken and grouped as Group1- silver nanoparticle,Group2-nanoparticle calcium hydroxide ,Group3- calcium hydroxide, Group4 – ciprofloxacin 20µg ( used only in MIC test). E.faecalis (MTCC 2729) is prepared according to 0.5 mac farland standard. Anti- bacterial activity and minimum inhibitory concentration is de- termined using disc diffusion method and micro broth dilution method respectively.

3. Methodology for antibacterial activity of silver nanoparticle, nanoparticle calcium hydroxide, calcium hydroxide on mature biofilm E. faecalis

3.1. Preparation of bacterial samplings

Teeth rinsed with sterile saline and wiped with alcohol outside. Sterile cotton pellets placed into chamber and access cavity is sealed with cavit.Apical foramen is sealed with hot glue to resemble closed system. Tooth is disinfected with 30% H2O2. Tooth surface coated with 10% tincture iodine and the sur- face is swabbed with 5% sodium thiosulphate.

3.2. Initial sampling

The Cavit and cotton pellets are removed and canal flushed with 2ml sterile saline and dried with sterile paper points. Sterile BHI broth inserted in to canals and removed with sterile paper points. This paper points are then placed into half of each BHI agar plates. Sterile BHI broth inserted and removed paper points are placed in 1ml test tube of BHI broth. The 1ml test tube is vortex- es and aliquot of 0.1 ml suspension was plated on to remaining half of BHI plates. The plates were incubated aerobically for 48 hours at 37°C.

3.3. Intracanal medicament placement

Irrigation with 20 ml of 1.5% NAOCL and 20 ml of sterile saline with side vented needles and Canal dried .Silver nanoparticle powder ( average particle size 20-30 nm, purity -99%, concentra- tion 1000ppm) is mixed with distilled water to paste like consist- ency and placed inside the canal using lentulo spirals ( group 1 – 8 teeth).Nano particle calcium hydroxide ( average particle size 30-50 nm, purity 99.5%) is mixed with distilled water to paste like consistency and placed inside the canal using lentulo spirals ( group 2 – 8 teeth ).Full strength calcium hydroxide ( ultra cal XS) is placed inside the canal ( group 3- 8 teeth).Access cavity were closed with 4mm Cavit and incubated at 37°C for 7 days. After 7 days canals are irrigated with 20 ml of 17% EDTA with side vented needles and rinsed with saline and dried with paper points. Group 4 – 8 teeth are negative controls, teeth with sterile incubations without medications.

3.4. Final sampling

Root canals of each tooth dried with sterile paper points and filled with reduced BHI broth. Three consecutive sterile paper points inserted in to the canal and paper points transferred to BHI agar plates. After collection of the first sample canals are filed with # 10k files and instrumented for 10 seconds. Dentinal shavings from files transferred to test tube containing 1ml of BHI broth. The test tube containing dentinal shavings is vortexed for 10 seconds, aliquots of 0.1 ml suspension were then placed on to another half of BHI agar plates and incubated aerobi- cally at 37 o C for 48 hours. CFU counted.

4. Results

All the materials are first subjected to antimicrobial test against E.faecalis by disc diffusion method. Ciprofloxacin 20 µg is used as positive control for this test. In this test group I AGNPs showed 11mm zone of inhibition at 500µg and Group II NCH showed 14mm zone of inhibition at 500µg. Group III CH do not exhibit any antimicrobial activity at this concentration. This clearly shows calcium hydroxide do not exhibit any antimicrobi- al property at very lower concentrations. The group IV positive control (ciprofloxacin 20µg) shows an average zone of inhibition of 35 mm at 500µg. Thereby this test clearly shows silver Nano- particles and Nanoparticle calcium hydroxide exhibit antibacte- rial efficacy against E. faecalis at a concentration greater than or equal to 500µg.Since calcium hydroxide did not exhibit any antimi- crobial activity at lower concentration it is excluded in the test for Minimum Inhibitory Concentration. MIC test was per- formed for AGNPs and NCH using serial dilution method. The test results are shown as optical density values and percentage of
inhibition. The test is repeated three times at every concentration in serial dilution and the average value is calculated for both the material. AGNPs shows an average optical density value of 1.082 and the percentage of inhibition is 37.77% at 25µg/200µl. NCH shows an average optical density value of 0.797 and percentage of inhibition is 54% at 12.5µg/200µl. The control group using ciprofloxacin showed optical density value as 1.738. Thus MIC for AGNPs is 25µg/200µl and NCH is 12.5µg/200 µl. The reduction in CFU before and after the placement of intracanal medicament is depicted in the following bar diagram.

The Mean CFU for group I before and after the placement of intracanal medicament is 5.74X10^6/ml and 1.37X10^7/ml respectively and showed 76% reduction in culture formation. Similarly the mean CFU/ml for group II before and after 7 days of its placement are 4.67X10^6 and 1.81X10^7 respectively and showed 61% reduction in culture formation. Group III shows mean CFU/ml before and after 7 days of its placement are 6.35X10^6 and 5.99X10^5 respectively and showed 11% reduction in culture formation. There is an increase in mean CFU/ml in group IV respectively and showed 11% reduction in culture formation followed by group II 61% and group III 11%.

### Table 1: Showing Student T-Test among Groups

<table>
<thead>
<tr>
<th>Groups</th>
<th>T-Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Agnp</td>
<td>14.3</td>
</tr>
<tr>
<td>Nch</td>
<td>8.34</td>
</tr>
<tr>
<td>Ch</td>
<td>1.54</td>
</tr>
<tr>
<td>Control</td>
<td>0.294</td>
</tr>
</tbody>
</table>

5. Discussion

The method used in the present study simulates the clinical conditions in teeth with infected root canals, in which microorganisms may lodge in areas of difficult access for the root canal preparation and medicaments (Menezes et al.,2004) [5]. E.faecalis was chosen because of its resistance to calcium hydroxide based intracanal medicaments.( Heling et al.,1992, Barrassi et al., 2003).This microorganism has been used in several studies(Evans et al.,2003, Siren et al.,2004, Delgado et al.,2010, Kandasswamy et al., 2010). [6 – 8]. The 21 day incubation period allowed the suspension of microorganism to diffuse throughout the root canal system (Heling et al, 1992). The CFU is counted after 7 days of intracanal medicament placement to verify the presence of viable bacteria. This time frame is chosen based on the previous study by Menezes et al., 2004, Cardoso et al., 2008. According to sjogren et al., 1991, a minimum contact time of 7 days is necessary for calcium hydroxide to eliminate microorganism that may have survived cleaning and shaping procedures. [3]

The antimicrobial activity of calcium hydroxide depends on the amount of free OH- ions released and its contact time with microorganism in the root canal. The carriers used as vehicle for calcium hydroxide decreases its PH and affect its antimicrobial activity. Water must be present for the antimicrobial effect of calcium hydroxide (Safavi k et al., 2000). [9] Calcium hydroxide in water has a thixotropic behavior and can be deposited in the canal with continuous agitation using lentulo spirals. Considering these facts, all the medicaments in this study are mixed to a paste like consistency and coated in the canal using lentulospirals. [1], [3]. The result of group I in this study correlates with the study done by Wu et al and kishen et al. [10] According to their results, sryringe irrigation with 0.1% AGNPs solution did not damage the biofilm structure. On the other hand, biofilms treated with 0.02% AGNPs gel as an intracanal medicament for 7 days significantly eradicated the structure with least number of residual viable E.faecalis cells in comparison with 0.01% AGNPs gel and calcium hydroxide groups.

The authors concluded that AGNPs should be applied as a medicament and not as an irrigant to exhibit potential activity against residual bacterial biofilm during root canal disinfection. [10] It is because the effectiveness of silver nanoparticle is directly proportional to the time it is in contact with bacterial cells. The probable mechanism of action of silver nanoparticle is the electrostatic attraction between the negative charge of the bacterial cell membrane and positively charged Nanoparticles (Kim et al., 2007). In terms of the molecular mechanisms of inhibitory action of silver ions on microorganisms, it has been shown that DNA loses its ability to replicate ( Feng et al.,2000) and the expression of ribosomal subunit proteins and other cellular proteins and enzymes necessary for ATP production becomes inactivated (Yamanaka et al., 2005). It has been hypothesized that silver ions affect membrane-bound respiratory enzymes (Bragg and Rainnie., 1974). The bacterial cells in contact with the silver Nanoparticles will take silver ions, which is possibly in turn will inhibit respiratory enzymes, and so help to facilitate reactive oxygen species/ free radical generation and subsequent damage to the cell membrane (Kim et al., 2007). [11] Similarly the results of group II in this study correlates with the study done by Omid Dianat et al [12] in which they concluded that the calcium hydroxide used in nanoparticle form as an intracanal medicament showed higher antibacterial efficacy than the calcium hydroxide and the nanoparticle calcium hydroxide had better tubular penetration thereby producing better antibacterial efficacy.

Intergroup comparison is done between following groups: Group I and Group II, the culture reduction is insignificant (p > 0.01). Similarly comparison in culture reduction between group III and group IV is also insignificant. All other combinations showed significant reduction in culture formation that is CFU/ml (p< 0.01).

Finally, Student- t - test is performed. Group I showed highest T value of 14.3, followed Group II 8.34, Group III 1.54 and Group IV showed the least value 0.294. The group I exhibit highest antibacterial efficacy among four groups followed by group II,III,IV. In our study GroupIII showed the least antibacterial efficacy. These results prove calcium hydroxide is inefficient in eliminating culture at a lower concentration in small time frame. In previous studies, it is reported that calcium hydroxide is a slow acting agent and should be kept in the canal for a longer period of time at a very high concentration,[14], [15] Most of the time it is difficult to achieve a high pH environment in root canal. It is mainly due to difficulty in depositing the intracanal medicament in the canal and poorer dentinal tubular penetration of the medicament. To overcome all these disadvantages, in this study calcium hydroxide has been used in Nanoparticle form (group II-NCH) and its antibacterial efficacy is compared with age known AGNPs (group I). And normal saline has been used as vehicle for Nanoparticles to avoid the disadvantages of vehicles interfering with its antibacterial efficacy due to its varied chemical
composition. And all the intracanal medicament used in this study are applied inside the canal using lentulospirals considering the thixotropic behavior of the mixtures for better tubular penetration. [13]

This study clearly demonstrates that the increased antibacterial efficacy of Nanoparticle is mainly due to its smaller size than their bulk or original counterparts thereby increasing its contact surface area and better tubular penetration.

6. Conclusion

From the results of the study it could be concluded that Nanoparticles had significant antimicrobial activity against Enterococcus faecalis as evidenced by the reduction. This preliminary study suggests that both the new nanoparticle antimicrobial agents tested here have the potential to be used as intracanal medicaments. But concerns such as duration of the medicament placement, hypersensitivity reactions, toxicity and effect on other organisms should be addressed in the future studies.

References