THE EFFECT OF THE ADDITION OF ACTIVATED CHARCOAL ON THE ANTIBACTERIAL EFFICIENCY OF DIFFERENT CALCIUM HYDROXIDE FORMULATIONS

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ABSTRACT

Aim of the study: Is to compare the antibacterial effect of conventional calcium hydroxide (GAMA dental lab, Cairo, Egypt), calcium hydroxide with Activated Charcoal (Eucarbon, Sedico, Cairo, Egypt), calcium hydroxide paste with iodoform (Metapex) (META,Chungcheongbuk-do,Korea) and calcium hydroxide paste with iodoform (Metapex) mixed with Activated Charcoal against *E-faecalis* using Agar diffusion method.

Materials and Methods: Samples were classified into 4 groups according to the material used. Each group was further divided into 3 subgroups according to the observation time (1 day, 3 days and 7 days). Each subgroup consisted of 7 agar plates implanted with *E-faecalis* strain. Four Holes were created in each agar plate each hole contained one tested material.

Results: Conventional Calcium hydroxide with saline had the largest inhibitory zone of *E-faecalis* in agar diffusion test over all observation period followed by Activated Charcoal and conventional calcium hydroxide followed by calcium hydroxide paste with iodoform with Activated Charcoal and the calcium hydroxide mixed with iodoform had no change in inhibitory zone.

Conclusion: The addition of activated charcoal did not improve the antibacterial effect of conventional calcium hydroxide while it improves the antibacterial efficacy of the calcium hydroxide and iodoform (Metapex).

INTRODUCTION

Chemical and biological dynamics of any intra canal medication is an important issue for choosing the desired intra canal medication. Microorganisms infecting the root canal system have a limited virulence when it is acting as an individual species however it is of high virulence when acting collectively. The success of endodontic treatment is highly dependent on the elimination of the infection from the root canal system, different intra canal medications has been used for this purpose however still the calcium hydroxide is considered the most popular and efficient intra canal medication used(1-5). So many

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changes were done to increase the antimicrobial action of calcium hydroxide that can speed up or slow down the ionic dissociation such as changing of the vehicle, addition of other active ingredients such as iodoform or changing of the particle size with use of the Nanotechnology, and adding different materials that may improve the antibacterial effect of calcium hydroxide formulations. The purpose of this study was to evaluate the effect of the addition of activated charcoal on the antibacterial efficiency of different calcium hydroxide formulations.

MATERIALS AND METHODS

Material

In this study, the following materials were used

a) Bacterial isolates

A total of (21) *E. faecalis* isolates were included in this study. All *E. faecalis* strains used in this study were isolated from human stool samples by culturing on MacConkey agar medium followed by subculture on nutrient agar medium for purification of the isolated colonies.

b) Substances:

- Calcium hydroxide Powder with saline.
- Calcium hydroxide with saline + Activated Charcoal conventional calcium hydroxide.
- Calcium hydroxide past with iodoform.
- Calcium hydroxide past with iodoform + Activated Charcoal.

c) Media:

The following media were used in this study:

1. Brain-Heart Infusion Broth:

   This medium was used to prepare the suspension of *E. faecalis*.

2. Brain-Heart Infusion Agar

   This medium was used to test the effect of the different substances on the growth of *E. faecalis* using the diffusion agar method.

Methods

a) Preparation of Brain-Heart Infusion Broth

   Thirty-seven grams of the medium were suspended in one liter of distilled water. Heating with frequent agitation ensured good mixing & dissolution. The suspension was then boiled for one minute until complete dissolution. It was later dispensed into appropriate containers and sterilized at 121°C for 15 minutes. The prepared medium was stored at 2-8°C. For best results, the medium was used on the same day.

b) Preparation of Brain-Heart Infusion Agar

   Fifteen grams of agar powder were added to 1 liter BHI broth & then heated to dissolve agar before dispensing into appropriate containers. Autoclaving was then performed for 15 min at 121°C to ensure adequate sterilization. The mixture was then poured into Petri dishes & left to cool & solidify.

c) Preparation of *E. faecalis* Suspension:

   A sterile swab was used to transfer bacterial growth from the primary culture into the BHI broth bottle & mixed well to form a homogenous suspension.

d) Classification of samples:

   21 samples were classified according to the tested material into 3 groups;

   - Group 1: Consisted of 7 holes filled with Calcium hydroxide Powder
   - Group 2: Consisted of 7 holes filled with Calcium hydroxide with saline + Activated Charcoal conventional calcium hydroxide
   - Group 3: Consisted of 7 holes filled with calcium hydroxide paste with iodoform
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Group 4: Consisted of 7 holes filled with calcium hydroxide paste with iodoform + Activated Charcoal. Each group was further divided into 3 subgroups according to the observation period to;

Subgroup A: One day.

Subgroup B: Three days.

Subgroup C: Seven days.

e) Implantation of the E. faecalis in the agar plates

A sterile cotton swab was dipped into the suspension and excess fluid was removed by turning the swab against the inside of the tube. The inoculum was evenly spread over the entire surface of dry BHI agar plates by swabbing in three different directions.

f) Mixing and application of the tested substances

Three holes of 4 mm diameter were made on the agar surface by a metal punch leaving about 10-15 mm away from the edge of the petri dish, and these holes were separated from each other by a distance not less than 20 mm to avoid overlapping zones of inhibition. Each hole contained one of the tested materials and marked as following;

- Hole number I for conventional Calcium hydroxide Powder.
- Hole number II for conventional Calcium hydroxide with Activated Charcoal.
- Hole number III for calcium hydroxide paste with iodoform and
- Hole number IV for calcium hydroxide paste with iodoform mixed with Activated Charcoal.

All the agar plates were incubated at 37°C in aerobic conditions for the required observation periods.

g) Method of evaluation

After one day observation the area of microbial growth inhibition (lack of bacterial colonization) around the holes were measured at the largest diameter with a poly gauge millimeter ruler. The agar plates were then re- incubated to take the measurements of the inhibitory zones after one day, three days and then after 7 days.

h) Statistical analysis

Data were analyzed by SPSS software (version 16.0, SPSS, Chicago, IL, USA). Data in each group were compared by the ANOVA and Kruskal-Wallis tests. Also the Dunnett’s test was performed to compare the results between two groups. The level of significance was set at 0.05.

RESULTS

A) Group one (Conventional Calcium hydroxide Powder)

The inhibitory zone after one day or 3 days or 7 days was the same changed to 22mm.

B) Group two (Conventional Calcium hydroxide Powder + Activated Charcoal)

The inhibitory zone after one day or 3 days or 7 days was the same changed to 21mm.

C) Group three (calcium hydroxide paste with iodoform)

There is no change of the inhibitory zone after one day or 3 days or 7 days.

D) Group four (calcium hydroxide past with iodoform + Activated Charcoal)

The inhibitory zone after one day or 3 days or 7 days was the same changed to 16mm.

DISCUSSION

The main purpose of endodontic treatment is the elimination of micobiota from the root canal system, which is the main challenge for all the endodontic practitioners, sporadic species of bacteria affecting the root canal system has low virulence impact.
however collectively they have high virulence impact.\textsuperscript{(9,10)} Estrela et al.\textsuperscript{(1)} Suggested the hypothesis of an irreversible inactivation of bacterial enzymes under extreme condition of PH for long period of time and also a temporary bacterial enzymatic inactivation with the restoration of normal activity when the PH returns to the ideal level. Estrela et al.\textsuperscript{(3)} Suggested the mechanism of action of calcium hydroxide through the release of hydroxyl ions that causing changes in the transport of nutrients and structure of organic component causing bacterial destruction. Lima et al.\textsuperscript{(11)} concluded that all calcium hydroxide medicaments were able to reduce colony forming unit (CFU) values of \textit{E-faecalis}. Chai et al.\textsuperscript{(12)} also concluded that calcium hydroxide are 100% effective in eliminating \textit{E- faecalis} biofilm.

Iodoform is composed of some powder with bright hexagonal crystals, it decomposes releasing iodine in nascent state.\textsuperscript{(13)} Compounds containing iodine are very employed for infection control in dentistry and it gives high radio-opacity for calcium hydroxide. The main problem of the action of the calcium hydroxide is that it has to be in direct contact with the bacteria to act on it, that’s why the addition of activated charcoal can be of benefit because of its adsorptive action that can attract the bacteria to the site of the calcium hydroxide and so can be beneficial to the antimicrobial effect of calcium hydroxide.

Agar diffusion test is a widely used test with reproducible results for the evaluation of the antimicrobial activity.\textsuperscript{(14)} It is able to demonstrate the activity of freshly mixed intra canal medications however it has many limitations as it lacks the ability to test and compare the viability of microorganisms used and the inability to distinguish between the bactericidal and bacteriostatic ability of tested materials.\textsuperscript{(15)} The results showed that with conventional calcium hydroxide, the addition of activated charcoal did not improve the inhibitory zone which may be related to the high spread of conventional calcium hydroxide powder mixed with saline. However with the use of calcium hydroxide with iodoform due to its putty consistency it has low spreading action through agar media and so no inhibition zone was detected while with the addition of activated charcoal with its adsorption effect it can attract the bacterial cells to the site of existence of calcium hydroxide and so improves its antibacterial efficacy.

**CONCLUSION**

The addition of activated charcoal did not improve the antibacterial effect of conventional calcium hydroxide while it improves the antibacterial efficacy of the calcium hydroxide and iodoform (Metapex).

**REFERENCES**


