Formulation and characterization of chlorhexidine HCl nanoemulsion as a promising antibacterial root canal irrigant: in-vitro and ex-vivo studies

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Introduction and aim: Chlorhexidine Hydrochloride [Chx.HCl] has a broad-spectrum antibacterial effect, sustained action and low toxicity so it has been recommended as a potential root canal irrigant. The aim of this study was to improve the penetration ability, cleansing and antibacterial effect of Chx.HCl using a newly formulated Chx.HCl nanoemulsion and use it as root canal irrigant.

Methods: Chx.HCl nanoemulsions were prepared using two different oils; Oleic acid and Labrafil M1944CS, two surfactants; Tween 20 and Tween 80 and co-surfactant; Propylene Glycol. Pseudoternary phase diagrams were constructed to designate the optimum systems. The prepared nanoemulsion formulae were evaluated for their drug content, emulsification time, droplet size, in-vitro drug release, thermodynamic stability, In-vitro antibacterial activity and ex-vivo study for the selected formula. Comparisons were made of Chx.HCl nanoemulsion with two different concentrations 0.75% and 1.6% vs Chx.HCl normal particle size as root canal irrigant for their penetration ability, cleansing effect and antibacterial effect.

Results: The selected formula was F6 with composition of 2% Labrafil, 12% Tween 80 and 6% Propylene glycol. It has small particle size (12.18 nm), short emulsification time (1.67 seconds), and fast dissolution rate after 2 minutes. It was found to be a thermodynamically/physically stable system. The higher concentration of Chx.HCl nanoemulsion 1.6% shows the best penetration ability compared to Chx.HCl normal particle size due to the smaller particle size. Chx.HCl nanoemulsion 1.6% has the lowest mean value of the remaining debris surface area (2001.47 µm²) when compared to normal particle size material (2609.56 µm²).

Conclusion: Chx.HCl nanoemulsion preparation has better cleansing ability and antibacterial effect with high efficacy on Enterococcus faecalis, where high reduction rate or complete eradication of bacterial cells has been achieved.

Keywords: chlorhexidine hydrochloride, nanoemulsion, root canal irrigant, penetration ability, cleansing effect, antibacterial irrigant

Introduction

Nanomulsions are a class of emulsions with a droplet size in the range of 50–500 nm and have attracted a great deal of attention in recent years because of their unique characteristics. 1 The growing use of non-ionic surfactants in nanoemulsions is related to their good detergent performance, which is not affected by water hardness, and, in most cases, to their low toxicity and they do not give rise to electrostatic interactions. 2 Nanotechnology with the ultra-small particle sizes, large surface area/mass ratio and unique physicochemical properties compared with their
bulk counterparts hold new prospects for the treatment and prevention of dental infections.\textsuperscript{3} Chlorhexidine hydrochloride (Chx.HCl) is sparingly soluble in water very slightly soluble in alcohol, it is gradually colored by light.\textsuperscript{4,5} Chx.HCl has a broad-spectrum antibacterial effect, sustained action and low toxicity. Because of these properties, it has also been recommended as a potential root canal irrigant. The major advantages of Chx.HCl are its lower cytotoxicity and lack of foul smell and bad taste.\textsuperscript{6–9} Several types of laser have been used to improve root canal disinfection. Bactericidal action of the laser depends on the wavelength and energy, and thermal effects that produce alteration of the cell wall of the bacteria, leading to changes in osmotic gradients up to cell death. The interaction between laser and the irrigant in the root canal outlines a new area of interest in the field of endodontic disinfection.\textsuperscript{10} Ultrasonic energy produces high frequencies but low amplitudes. The files are designed to oscillate at ultrasonic frequencies of 25–30 kHz, which are beyond the limit of human auditory perception (>20 kHz). They operate in a transverse vibration, setting up a characteristic pattern of nodes and antinodes along their length. The term passive ultrasonic irrigation (PUI) is an irrigation scenario where there is no instrumentation or contact of the canal walls with an endodontic irrigant. During PUI, the energy is transmitted from the irrigant to the root canal by means of ultrasonic waves. The latter induces acoustic streaming and cavitation of the irrigant.\textsuperscript{11} Based upon the above evidence, it was thought that evaluation by using nanotechnology to improve the penetration ability and cleansing effect of Chx.HCl is of value.

**Materials and methods**

**Materials**

Chlorhexidine hydrochloride Chx.HCl was kindly provided by the Arab Drug Company for Pharmaceuticals (Cairo, Egypt). Labrafil M 1944 CS (oleoyl polyoxyyl-6 glycerides), was generously supplied by Gattefosse (Saint-Priest, France). Tween 20 (polyoxyethylene (20) sorbitanmonolaurate), Tween 80 (polyoxyethylene (80) sorbitanmonooleate), Oleic acid, Propylene glycol were supplied by Gomhorya Company (Cairo, Egypt). Noncarious single rooted human teeth were extracted due to periodontal diseases or for orthodontic treatment in the oral and maxillofacial department in the Faculty of Dentistry, Ain Shams University, Cairo, Egypt. Pure culture of *E. faecalis* (strain ATCC 29212) grown in brain heart infusion (BHI) broth (RC CLEANER, Ichung Dental Ltd., Seoul, Korea).

**Methods**

**Solubility studies**

Solubility of Chx.HCl in different media (Oleic acid, Labrafil M 1944CS, Tween 20, Tween 80, Propylene glycol and water) were studied. Large excess of Chx.HCl (50 mg) was taken in centrifuge tubes and 5.0 g media phase was added. The mixture was shaken in a vortex mixer for 15 minutes and then stored at room temperature. After 24 hours, in order to obtain clear supernatant, the undissolved drug precipitates in the tubes were centrifuged for 5 minutes at 3000 rpm. Adequate sample solution was collected and diluted with n-butanol. Whatman 102 filter paper was used to filter the diluted sample which was further suitably diluted with n-butanol to determine the concentration of drug in saturated solution. The samples were analyzed with UV spectrophotometer at 260 nm against n-butanol as blank.\textsuperscript{12,13}

**Construction of a pseudoternary phase diagram**

A pseudoternary phase diagram was constructed to determine the exact proportion of each component needed in the formulation to obtain the best parameters for the ideal nanoemulsion.\textsuperscript{14} Formulations were prepared using oil, namely Oleic acid and Labrafil M1944CS, surfactants, namely Tween 20 and Tween 80 and co-surfactant, namely Propylene glycol. Firstly, a mixture of surfactant alone (with no co-surfactant) and oil was prepared in different volume ratios (from 1:9 to 9:1). When aqueous titration (dropwise addition of water) was performed on the mixture, the observation of clear to turbid mixture, which was the end point, was closely monitored. These end points were then marked on a pseudoternary phase diagram. The whole procedure was repeated for a mixture of surfactant and co-surfactant \( S_{\text{mix}} \) which were prepared in ratios 2:1 and 3:1 and combined with the selected oil.\textsuperscript{15,16} Figure 1.

**Preparation of Chx.HCl nanoemulsion**

Nanoemulsion systems containing Chx.HCl were prepared using Labrafil M 1944 CS as oil phase and surfactants Tween 80 or 20 as well as, Propylene glycol as cosurfactant, and, lastly, water, Table 1. The drug was dissolved in the Labrafil M 1944 CS, the water combined with surfactant and a cosurfactant is then added at a slow rate with gradual stirring. The amount of surfactant and cosurfactant to be added and the percent of oil phase that can be incorporated
were determined with the help of pseudo-ternary phase diagram. Ultrasonicator (Ultrasonic LC 60 H, Elma, Germany) was used to achieve the desired size range for dispersed globules. It was then allowed to equilibrate.\(^\text{17}\)

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**Characterization of Chx.HCL nanoemulsion**

**Dispersibility tests**

Dispensibility tests were done using a dissolution apparatus (Dr. Schleuniger Pharmaton, type Diss 6000, Thun,
Switzerland) where 1 mL of each formula was added to 500 mL of water at 37±0.5°C. A standard stainless-steel dissolution paddle rotating at 50 rpm provided gentle agitation. The type of emulsion formed was visually inspected and categorized as either clear, translucent with bluish tone, milky or turbid emulsion. The formulae that have clear appearance were selected for further studies.18,19

Determination of drug content

Chx.HCl from an optimized nanoemulsion formula based on the pseudoternary phase diagram results was extracted in n-butanol using the sonication technique. The extract was analyzed for Chx.HCl content spectrophotometrically at a wavelength of 260 nm after suitable dilution.20

Test for self-emulsification time of Chx.HCL nanoemulsion

To test the self-emulsification time, 1 mL volume of each formula was introduced into a beaker containing 250 mL of distilled water, maintained at 37±1°C under continuous stirring at 50 rpm. The self-emulsification time was taken as the time for a pre-concentrate to form a homogenous mixture upon dilution.21

Droplet size analysis

For the droplet size analysis 50 mg of the optimized formula were diluted with water to 1000 mL in a flask and gently mixed by hand. The droplet size distribution was determined by Malvern Zetasizer 2000 (Malvern Instruments Ltd., Malvern, UK) with the conditions of backscatter detection at 173°; temperature of 25°C; refractive index of 1.330 were used.22

In-vitro release study

The in-vitro dissolution studies were carried out using USP apparatus type II (paddle) (Dr. Schleuniger Pharmaton, type Diss 6000) at 50 rpm. The dissolution medium used was distilled water (500 mL) maintained at 37±0.5°C, and 5 mL of prepared formula was instilled to the dissolution medium. Then 5 mL of dissolution media were withdrawn at different time intervals and the content of drug released was determined spectrophotometrically at 254 nm. The experiments were conducted in triplicate.23

Kinetic study of the release data of ChxHCL nanoemulsions

The kinetic parameters of the in-vitro release of Chx.HCl from its prepared nanoemulsions were then measured. Zero, first and second order kinetics and the Higuchi-diffusion model were tried to choose the most suitable kinetic order for Chx.HCl release.

Thermodynamic stability tests of prepared Chx.HCl nanoemulsions

Phase separation and drug precipitation

A quantity of 2 mL of each formula was stored for 48 hours at ambient temperature and observed thereafter for phase separation. Then 1 mL samples of each Chx.HCl nanoemulsion formula was diluted to 10 mL and 100 mL with distilled water at 25°C stored for a period of 24 hours. and observed afterwards for phase separation.21

Refrigeration thaw cycle

Next, 2 mL samples of each formula were separately transferred to a transparent screw capped bottle and stored in a refrigerator at 2°C for 24 hours. After which they were removed and stored at 25°C and 40°C. A single refrigeration thaw cycle test was performed. The samples were then observed for phase separation and drug precipitation.21

Centrifugation

A 5 mL sample of each Chx.Hcl nanoemulsion formula formulation was transferred into a glass test tube and inserted into a laboratory centrifuge (Microcentrifuge Shanghai Surgical Instrument Factory, model 800, Shanghai, People's Republic of China) and centrifuged at 4,000 rpm for 5 min. Thereafter, the samples were observed for phase separation and drug precipitation.21

Irrigating solution examinations on extracted teeth

Penetration ability and cleansing effect

All experiments were approved by the Institutional Ethics Committee, Ain Shams University, Egypt. Fifty noncarious single rooted human teeth with mature apices were selected. The extracted teeth were used after taking signed written informed consent letters from patients. The teeth included maxillary and mandibular incisors and mandibular premolars. The external root surfaces were debrided using a curette and all teeth were placed in 0.5% NaOCl for 24 hours for surface disinfection and then stored in sterile saline until use. The crowns were removed using a safe sided diamond disc and the tooth length was standardized to 16 mm from the root apex to the coronal border.24,25 The teeth were divided into the following groups according to the solution of irrigation:
Group (A) (n=24) the samples were irrigated with Chx. HCl nanoemulsion. Subgroup (I) (n=12) the samples were irrigated with 5 mL Chx.HCl nanoemulsion 0.75% concentration. Subgroup (II) (n=12) the samples were irrigated with 5 mL Chx.HCl nanoemulsion 1.6% concentration. Group (B) (n=24) the samples will be irrigated with 5 ml 2% Chx.HCl normal particle size. Control groups: (n=2), irrigation with 5 mL saline No activation.

Each subgroup of A and B was divided according to the method of activation of the irrigant:Laser

(a) (n=4): Laser treatment was performed with a diode laser (Epic x diode laser 940, Biolase, Irvine, CA, USA) and the endodontic tip at a wavelength of 940 nm and output power of 1.5 w according to the instructions of the manufacturer. The laser irradiation was delivered for 30 seconds into the canal to the working length.

(b) Passive ultrasonic irrigation (PUI) (n=4): The ultrasonic tip was inserted in each canal full of irrigant for 30 seconds (Satelec P5, Acteon® Group, Merignac, France, Irrigation tip size 25).

(c) No activation (n=4): Mechanical preparation was done using standardized technique to a master apical file size 50. All the samples were grooved on the mesiodistal surfaces along their entire lengths and split longitudinally in two halves using a mallet and chisel. Then all samples were examined under scanning electron microscope (SEM).

Antibacterial effect
Forty-four noncarious single-rooted human teeth with mature apices were selected. The teeth included maxillary and mandibular incisors and mandibular premolars. The external root surfaces were debrided using a curette and all teeth were placed in 0.5% NaOCl for 24 hours for surface disinfection and then stored in sterile saline until use. The crowns were removed using a safe sided diamond disc and the tooth length was standardized to 16 mm from the root apex to the coronal border.

Mechanical preparation was done using standardized technique to a master apical file size 50. Sterile saline was used as irrigant during the procedure. The canals were finally flushed with 2 mL 17% EDTA for 1 min to remove the smear layer. The entire root surfaces, including the apical foramen of each sample were coated with two layers of nail polish (cyanoacrylate adhesive) to prevent leakage. The teeth were then mounted vertically in dental stone blocks to make handling and identification easier. The samples were then autoclaved at 121°C and 15 lb pressure for 20 minutes. Subsequent to sterilization all samples were transported and manipulated under aseptic conditions using sterile instruments. Pure culture of E. faecalis (strain ATCC 29212) grown in brain heart infusion (BHI) broth for 24 hours at 37°C was used to contaminate the root canals. An inoculum of pure suspension of E. faecalis was suspended into the prepared canals of all teeth by using sterile micropipette. The blocks were then placed inside sterile beakers and incubated at 37°C for 24 hours.

The teeth were divided into the following groups according to the solution of irrigation:

Group (A) (n=24) the samples were irrigated with Chx. HCl nanoemulsion. Subgroup (I) (n=12) the samples were irrigated with 5 mL Chx.HCl nanoemulsion 0.75% conc. Subgroup (II) (n=12) the samples were irrigated with 5 mL Chx.HCl nanoemulsion 1.6% conc.

Group (B) (n=12) the samples will be irrigated with 5 mL 2% Chx.HCl normal particle size.

Control Groups: Positive Control, (n=4) the contaminated root canals were irrigated with 5 mL normal saline and kept as positive control. Negative Control: (n=4) specimens did not receive suspension injections, ie, root canals were not contaminated with E. faecalis and were kept sterile to serve as negative controls to confirm sterilization and the reliability of the procedures. A volume of 5 mL of the test irrigant was used in each sample. Then final irrigation was done with 1 mL sterile normal saline for each sample.

Each subgroup of A and B was divided according to the method of activation of the irrigant:

(a) Laser (n=4), Laser treatment was performed with a diode laser (Epic x diode laser 940, Biolase) and the endodontic tip at a wavelength of 940 nm and output power of 1.5 w according to the instructions of the manufacturer. The laser irradiation was delivered for 30 seconds into the canal to the working length.

(b) Passive ultrasonic irrigation (PUI) (n=4), the ultrasonic tip was inserted in each canal full of irrigant for 30 seconds (Satelec P5, Irrigation tip size 25).

(c) No activation (n=4).

Evaluation of bacterial growth
Sterile paper point size 35 was used to collect samples from root canals. The paper point was inserted into the
canals till the working length, left for 10 seconds and then transferred to agar plates to determine the number of colony-forming units (CFUs) per plate. The plates were incubated at 37°C for 24 hours after which the plates were visually evaluated for occurrence of bacterial growth. Clear plates demonstrated complete sterilization. Blurred plates were considered showing positive growth. The mean number of CFUs in the areas of bacterial growth on each plate was determined and the number of CFUs was calculated. Survivors were measured primarily using viable counts on drop plates. Additionally, pour plates were used to count low colony-forming units and dilutions to 106 were used to count high CFU.\(^{36,37}\)

**Viable count method using the pour plate method\(^{38}\)**

A tube containing 15 mL of melted agar medium was prepared where the agar medium was pre-sterilized in an autoclave on the same day it was needed for the experiment. *E. faecalis* a facultative Gram positive anaerobic coccus has the capability of surviving at very high pH, acidity and high temperatures.\(^{39}\)The bacterial sample (*E. faecalis* ATCC 29212) was prepared by mixing cells from a colony into sterile saline solution. Then bacterial sample was diluted with saline to match McFarland 0.5 which is equivalent to 108 cfu/mL. The sample volume added was 10 µL.\(^{39}\) Preparation of Turbidity standard (McFarland 0.5)\(^{40}\) was achieved by pouring 0.6 mL of a 1% (10 g/L) solution of barium chloride dihydrate into a 100 mL graduated cylinder, and filling to 100 mL with 1% (10 mL/L) sulfuric acid. The turbidity standard was placed in a tube identical to the one used for the broth sample and stored in the dark at room temperature for 6 months and sealed well to prevent evaporation. The lid of the empty Petri dish was opened, and the sample was dispensed into the middle of the plate. If the agar had thoroughly solidified then the plate was inverted and incubated at 37°C for 24 hours.

**Statistical methods**

All data were collected, tabulated and subjected to statistical analysis. Statistical analysis was performed with IBM\(^{®}\) SPSS\(^{®}\) Statistical Version 17 for Windows (SPSS Inc., IBM Corporation, Armonk, NY, USA).

**Results and discussion**

**Solubility studies**

The solubility of Chx.HCl in various oily phases, surfactant solutions, cosurfactant solution, and water were studied. The highest solubility of Chx.HCl was in Labrafilm M and the least one was in Oleic acid. The higher solubility of the drug in the oil phase is important for the nanoemulsion as the ability of the nanoemulsion to maintain the drug in solubilized form means that higher solubility of the drug in oil will ensure a lesser amount of oil in the formulation and consequently lesser amounts of surfactants and co-surfactants are required for emulsification of drug loaded oil droplets.

**Construction of a pseudoternary phase diagram**

A pseudoternary phase diagram was constructed to identify the nanoemulsifying region and to optimize the concentration of the selected oil, surfactant, and co-surfactants (namely Labrafilm M, Tween 80, Tween 20 and propylene glycol, respectively). Chx.HCl showed very low solubility in oleic acid which resulted in a turbid appearance when titrating Oleic acid with water from the first drop. So oleic acid systems were excluded from this study. Other prepared formulae gave clear solutions with an infinite amount of water during titration when prepared with an oil and surfactant mixture in a ratio of 1:9. The surfactants used (Tween 20 and Tween 80) are non-ionic hydrophilic surfactants and were selected due to lower oral toxicity with better stability to emulsion over a wide range of pH and ionic strength.

**Characterization of Chx.HCL nanoemulsion**

**Dispersibility tests**

All prepared formulae showed a clear appearance except system F2 which showed a turbid appearance, so it was excluded from further evaluation study.

**Determination of drug content**

The drug content of the prepared Chx.HCl nanoemulsion formulae ranged between 86.3 and 96.7%.

**Test for the self-emulsification time of Chx.HCL nanoemulsion**

An ideal nanoemulsion formulation should have the ability to disperse completely and quickly when subjected to dilution under mild agitation. Chx.HCl nanoemulsion formulae showed short emulsification time ranged between 1.67 and 12.33 seconds. Tween 80 was found to have the least emulsification time. This could be explained by the higher solubilization capacity of Tween 80. Self emulsification time was increased by the increase in the concentrations of surfactant which may be...
due to the increase in the viscosity of the system by the surfactant.

**Droplet size analysis**

The droplet size of the emulsion defines the rate and extent of drug release. Smaller emulsion droplet size results in lower emulsification time and a larger surface area provided for the drug absorption. The average droplet size of selected Chx.HCl nanoemulsion formulae were 711±0.44, 587±15.3, 10.97±0.11, 16.43±4.55 and 12.18±2.48 with PDI 0.76, 0.19, 0.61, 0.47, and 0.16 for F1, F2, F3, F4, F5, and F6 respectively. The formulation containing Tween 80 as surfactant showed smaller globule size. This might be attributed to its higher emulsifying ability. PDI values were low, representing that the system had a narrow size distribution. These formulae showed clear appearance as their droplet radii fell below the optical wavelength of visible light (390–750 nm) by which minimum light scattering took place.

**In-vitro release study**

Figure 2 shows the percent of Chx.HCl release from the prepared formulae. The complete drug release for prepared Chx.HCl nanoemulsion formulae ranged between 2 and 7 mins. It was observed that the highest drug release rate was obtained in the case of Chx.HCl nanoemulsion formula F6 (2 minutes) and that may be attributed to the presence of Tween 80 which exhibits higher emulsifying, the resulting nano-level droplet size which provided a large surface area for the release of the drug, thus permitting a faster rate of drug release. In the same time the solubility property of propylene glycol allowed large quantities of the hydrophilic surfactant to dissolve in the oil.

**Kinetic study of the release data of Chx. HCL nanoemulsion**

It was found that the in-vitro release of Chx.HCl followed different kinetic orders and no definite kinetic order could express the drug release from differently prepared nanoemulsion formulae. The kinetic release of drug from F4 was first order kinetics, which indicates that they release in a way that is proportional to the amount of drug remaining in its interior. The kinetic release of drugs from other formulae followed the Higuashi Diffusion model which indicates that the amount of drug release is proportional to the square root of the total amount of drug and the solubility of the drug in the nanoemulsion formula.

**Thermodynamic stability tests of prepared Chx.HCL nanoemulsion**

The selected formulae were subjected to different thermodynamic stability by using heating cooling cycles, centrifugation and freeze-thaw cycle stress tests. It was observed that F3 and F4 formulae exhibited drug precipitation after a thaw cycle, and F1 exhibited thickening (gelation). F5 and F6 have passed the successive cycles of centrifugation, heating-cooling test and freezing-thawing test. Nanoemulsions are thermodynamically stable systems and are formed at a particular concentration of oil, surfactant, and water, with no phase separation, creaming or cracking. It is the thermo-stability which differentiates nanoemulsions from emulsions that have kinetic stability and will eventually phase separate. F3 exhibited larger particle size than other formulae (587 nm) which may be the cause of phase separation and drug precipitation in the thermodynamic stability test. F4 which includes Tween 80
with no cosurfactant exhibited drug precipitation which may indicate the need for using propylene glycol with Tween 80 for nanoemulsion formula stability. F1 which includes Tween 20 with no cosurfactant exhibited thickening (gelation) which is an increase in the viscosity or gel strength due to droplet aggregation.

The stability results confirm the importance of presence of the cosurfactant propylene glycol to increase dispersion of particles and prevent drug precipitation.\(^{43}\) F6 was the best formula as it has small particle size (12.18 nm), short emulsification time (1.67 seconds), and fast dissolution rate after 2 minutes. It was found to be a thermodynamically/physically stable system, so it was selected for further study.

**Irrigating solution examinations on extracted teeth**

**Penetration ability in dentinal tubules**

Failures after root canal treatment are increasingly frequent representing elevated risk of more complex infections in patients.\(^{44,45}\) The removal of the biofilm is mandatory in root canal disinfection and sealing.\(^{46,47}\) Due to the complex nature of the root canal system it becomes difficult for the complete debridement of canal containing bacteria with instrumentation and irrigation alone.\(^{48}\) Effectiveness of an endodontic irrigating solution is dependent on the penetration of irrigant in DT and duration of action against bacteria.\(^{49}\) Hence, newer modalities were tried and tested for thorough disinfection of root canals. Conventional irrigants cannot completely eliminate *E. faecalis*, because of lesser penetration depth into DT.\(^{50}\)

**Effect of irrigant type**

The mean of irrigant cleansing ability for nanoemulsion irrigants was 2001.47 µm\(^2\) while for normal particle size irrigants the mean was 2609.56 µm. The mean difference between nanoemulsion irrigants and normal particle size irrigants was 608.09 µm\(^2\). There was a statistically highly significant (\(P<0.001\)) difference between nanoemulsion irrigants and normal particle size irrigants with (\(P\)-value 0.00052). Nanoemulsion had a statistically highly significant difference showing a lower mean value of the remaining debris surface area when compared to normal particle size material, ie nanoemulsion materials had better cleansing ability as shown in Figure 3.

**Effect of irrigants’ concentration**

The mean surface area of remaining debris for Chx.HCl 1.6% was 2320.36 µm\(^2\) when compared to Chx.HCl 2% its mean was 2949.85 µm\(^2\). There was a statistically highly significant (\(P<0.001\)) difference between the higher concentration of nanoemulsion irrigants and the normal particle size irrigants (\(P\)-value 0.00000). Although the concentration of nanoemulsion irrigants was lower in comparison to the concentration of normal particle size irrigants, the effect of this low concentration on debris removal was significantly stronger and more effective in cleansing the canals.

**Effect of activation method**

PUI had a statistically highly significant difference (\(p<0.001\)) when compared to other activation methods. The mean value of remaining debris surface area was 1695.31 µm\(^2\) when activated by PUI. The mean difference between PUI and Laser was 987.89929 showing highly statistically significant (\(P<0.001\)) difference with (\(p\)-value 0.00000). The mean difference between PUI and no activation was 712.40643 showing a highly statistically significant (\(P<0.001\)) difference with a \(p\)-value of 0.00098). The use of either Laser activation or no activation was not significantly statistically (\(P>0.05\)) different with a \(P\)-value of 0.451211. The mean surface area of the remaining debris when Laser activation was used was 2683.21 µm\(^2\). The mean surface area of the remaining debris when no activation was used was 2407.72 µm\(^2\). PUI has a statistically lower mean value of debris surface area when compared to either laser activation or no activation, ie, better cleansing ability.

**Cleansing ability**

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Figure 3 Comparison between irrigants cleansing ability: (A) Nano CHX laser activation, (B) CHX laser activation, (C) Nano CHX PUI, (D) Nano CHX no activation, (E) CHX no activation and (F) CHX PUI activation.
significantly statistical (P>0.05) different with (P-value 0.451211). The mean surface area of remaining debris when Laser activation was used was 2683.21 µm². The mean surface area of remaining debris when no activation was used was 2407.72 µm². PUI has statistically lower mean value of debris surface area when compared to either laser activation or no activation, ie better cleansing ability.

Comparison between irrigants cleansing ability

The mean effect of nanoemulsion irrigants on debris removal were statistically significantly higher than normal particle size irrigants. Chx.HCl 1.6% with PUI had 1938.77 µm²; while with the laser it was 2510.96 µm². With no activation the mean was 2511.34 µm². The worst results, largest amount of debris, was seen when Chx.HCl 2% was used and activated by Laser. The same results were also seen when Chx.HCl 0.75% had no activation. It is clear that the best results were achieved when the higher concentration of the irrigant in nanoemulsion dosage form was used. PUI has the most potent effect in activating the irrigants and flushing debris out as shown in Figure 3A–F).

Antibacterial effect

Viable count method using the pour plate method

As shown in Table 2 nanoemulsion Chx.HCl had a better result than normal particle size in terms of viable count with good correlation with the formulae penetration ability and cleansing effect according to the following parameters: Irritant type either nanoemulsion or normal particle size, irrigant concentration, and methods of activation.

Complete bacterial elimination was achieved when the higher concentrations of irrigants were used. Chx.HCl 0.75% had the worst antibacterial effect even by activation with PUI. Laser activation had an adverse effect on the nanoemulsion irrigant. From all the previous results it was obvious that the use of laser has reduced the efficiency of Chx.HCl 0.75% nanoemulsion, where the CFU for nano Chx.HCl 0.75% was 195 which is very high count indicating that the reagents were not effective in this concentration with laser activation. The diode laser has photothermal effects so either the light or the heat may cause the nanoemulsion to lose its antibacterial effect. The results for the high concentrations were complete bacterial eradication. Nano Chx.HCl 1.6% showed negative bacterial growth with the presence of laser activation which means that laser did not affect the antibacterial ability of nano Chx.HCl 1.6%. It can be concluded that the higher concentration of nanoemulsion material has a better antibacterial effect.

Conclusion

In this work, Chx.HCl nanoemulsions were prepared using two different oils, two surfactants and co-surfactant; the optimum formula (F6) that displayed small particle size,

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<tr>
<th>Positive control</th>
<th>Bacteria only on Agar +++++ve</th>
<th>Bacteria only in the canals ++ve</th>
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<td><strong>Nanoemulsion</strong></td>
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<tr>
<td><strong>Material concentrations</strong></td>
<td>CHX I 0.75%</td>
<td>CHX II 1.6%</td>
</tr>
<tr>
<td><strong>Negative control</strong></td>
<td>-ve</td>
<td>-ve</td>
</tr>
<tr>
<td><strong>Activation methods showing number of viable colonies per plate</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1- Laser</td>
<td>195</td>
<td>-ve</td>
</tr>
<tr>
<td>Cfu/ml</td>
<td>19500</td>
<td>-ve</td>
</tr>
<tr>
<td>2- PUI</td>
<td>168</td>
<td>-ve</td>
</tr>
<tr>
<td>Cfu/ml</td>
<td>16800</td>
<td>-ve</td>
</tr>
<tr>
<td><strong>Without activation showing number of viable colonies per plate</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>180</td>
<td>95</td>
<td>120</td>
</tr>
<tr>
<td>Cfu/ml</td>
<td>18000</td>
<td>9500</td>
</tr>
</tbody>
</table>
short emulsification, and fast dissolution rate, was selected. Additionally, (F6) was subjected to thermodynamical/physical stability tests. In the Chx.HCl nanoemulsion in a concentration of 1.6% compared to traditional Chx.HCl as an irrigation solution, the Chx.HCl nanoemulsion shows the best penetrations in dentinal tubules, and cleansing ability with PUI as the activation methods. Moreover, the antibacterial study of Chx.HCl nanoemulsion shows complete bacterial elimination. The results confirmed that Chx.HCl nanoemulsion could be considered as a promising irrigant solution.

Acknowledgments
We are so grateful to the staff members of the research laboratory, Misr University for Science & Technology, for their kind support.

This research did not receive any specific grant from funding agencies in the public, commercial, or not-for-profit sectors.

Disclosure
The authors report no conflicts of interest in this work.

References


