Effectiveness of Cryogen Tetrfluoroethane on Elimination of Gingival Epithelium and its Clinical Application in Gingival Depigmentation–Histological Findings and Case Series

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ABSTRACT

Objective: To histologically assess and clinically co-relate the effectiveness of cryogen Tetrfluoroethane (TFE) for gingival depigmentation procedure.

Material and Methods: Twelve patients having unaesthetic gingival melanin pigmentation were included in the study. Gingival tissues of eight patients having gingival melanin pigmentation undergoing gingivoplasty or gingivectomy for crownlengthening were exposed to the cryogen and this was used for the histological examination. Gingivectomies were done after 8, 24, 96 hours and after a week of application of tetrfluoroethane. Four fair skinned patients complaining of unaesthetic gingival hyperpigmentation underwent gingival depigmentation using Tetrfluoroethane cryogen.

INTRODUCTION

Low temperature was commonly used to achieve specific effect on tissues and its popularity has increased from the past 3–4 decades [1]. Cryosurgery is an effective method of tissue destruction by freezing and has become firmly established surgical technique for treatment of diverse benign and malignant cutaneous lesions [2] and it is also used in dental practice [3].

In 1969, Heitmann, Dorman and Collins [4] reported that the gingiva freezes at a temperature ranging between 14.2°C to a high of 31.8°C may be conducted without adversely affecting the vitality of adjacent teeth. With a melting point of -101°C and a boiling point of -26°C, Tetrfluoroethane is commercially available in a pressurized spray can and immediately evaporates without residue following spraying. This gaseous fluorocarbon is used in dentistry in the field of endodontics for cold-pulp testing for full crowned or natural teeth and in orthodontic treatment to facilitate the seating of nickel-titanium expansion loops [5].

Tetrfluoroethane is safe, exposure via shortterm inhalation and skin contact show no effect on inhibition pulse, blood Pressure, electrocardiogram or lung function in healthy volunteer [6]. Tetrfluoroethane exerts no adverse effect on development, maturation and reproduction and does not have genotoxic and oncogenic capacity on animals [7-9].

The cryogen used in this study was obtained in a pressurised can and dispensed via outlet nozzle from the can. The present study attempted to histologically assess and clinically co-relate the effectiveness of ultralow temperature using Tetrfluoroethane in elimination of gingival epithelium in turn resulting in depigmentation.

RESULTS

Histologically after 96 hours of application of cryogen there was complete loss of rete pegs and epithelial detachment from the corium was evident. Complete re-epithelialisation was noted after a week and was clinically correlated.

CONCLUSION

We therefore, concluded that histologically tetrfluoroethane can effectively destroy gingival epithelium without causing damage to the connective tissue and clinically the color of the gingiva had more pleasing appearance 6 months postoperatively. Hence the cryogen can be used safely for depigmentation procedure.

MATERIAL AND METHODS

Twelve patients were used as experimental subjects for the study.

1. Clinically all were free of any form of gingivitis.
2. A detailed medical history which included pregnancy, breast feeding, systemic diseases which were associated or not associated with the gingival melanin pigmentation, medications and adverse drug reactions were taken into consideration. The relative contraindications included cold intolerance and cold urticaria.
3. Marginal gingiva including the interdental papillae of the maxillary cuspid and bicuspids of patients undergoing gingivoplasty or gingivectomy for crownlengthening and had melanin pigmentation were frozen.
4. A total of eight gingivae obtained from gingivectomy for crown lengthening were frozen.
5. Freezing procedure: a. The tissue to be frozen was isolated, dried; topical anaesthetic (10% Xylocaine, Andheri east, Mumbai, India) was sprayed.
   b. The cryogen was dispensed in a kidney tray and was transferred to the site using a cotton pledget carried using a tweezer. It was applied for 2-3 minutes without applying too much of pressure [Table/Fig–1A].
   c. Thawing occurred within 10-12 seconds after removal of the cotton pledget from the site.
6. Biopsies were taken in the following order: 8, 24, 96 hours and one week after freezing [Table/Fig–1B].
7. The patients undergoing cryosurgical depigmentation had generalized diffused pigmentation score of 3 classified according...
to the gingival pigmentation index (GPI) [10] as shown below. The patients were reviewed after 6 months of depigmentation.

<table>
<thead>
<tr>
<th>Score</th>
<th>Criteria</th>
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<tr>
<td>0</td>
<td>Absence of pigmentation, pink color of the gingiva</td>
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<tr>
<td>1</td>
<td>Spots of brown to black</td>
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<tr>
<td>2</td>
<td>Brown to black pigmentation, more than spots but not diffuse (patches of pigmentation)</td>
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<tr>
<td>3</td>
<td>Diffuse brown to black pigmentation involving papillary, marginal and attached gingiva</td>
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**RESULTS**

**Clinical signs**

Immediately after the application of the cryogen a solidly frozen area resembled frozen tissue. Thawing was evident and recognized as the size of the frozen area decreased and completely disappeared in 15-20 seconds. The appearance of site after thawing was indiscernible from adjacent site with slight redness.

After 8 hours the site showed shallow elevated surface and in 96 hours the overlying epithelium easily peeled off exposing the underlying connective tissue. After a week of application the overlying epithelium was completely replaced by a depigmented epithelium.

**Histological findings**

After 8 hours [Table/Fig-2A] on examination the para keratinized stratified squamous epithelium showed loss of rete pegs. There were some degenerative changes in the form of necrotic material. Basement membrane was inconspicuous. Underlying connective tissue stroma showed severe chronic inflammatory response mainly in the form of lymphocytes and plasma cells. After 24 hours histological examination of the specimen showed parakeratinised stratified squamous epithelium with acanthosis at places. The epithelial cells show desquamative changes in the form of hydropic degeneration [Table/Fig-2B]. Underlying connective tissue stroma is loose and oedematous with few chronic inflammatory cells in the form of lymphocytes and plasma cells.

After 96 hours the soft tissue showed parakeratinised squamous epithelium exhibiting degeneration. Underlying connective tissue is delicate to dense with focal aggregate of chronic inflammatory cells [Table/Fig-3A].

Clinically the gingival tissue surface showed desquamation and redness suggesting healing and replacement with a lighter colored gingiva [Table/Fig-3B] the pigmentation of the gingiva changed to a score of 0 in three out of four cases and score of 1 in one case.

After a week of application the cryogen there was complete reepithelialisation [Table/Fig-4A] and formation of rete pegs with the gingival pigmentation index (GPI) [10].

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<td>0</td>
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<td>1</td>
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**Table/Fig-1A**: Application of the cryogen

**Table/Fig-1B**: Excised tissue sent for histological examination

**Table/Fig-2A**: Eight hours following freezing showing epithelial degeneration

**Table/Fig-2B**: Specimen after 24 hours showing loss of rete pegs

**Table/Fig-3A**: Four day specimen showing complete loss of rete pegs and subepithelial cleft

**Table/Fig-3B**: Clinical resemblance after four days of application

**Table/Fig-4A**: Specimen after a week, showing complete reepithelialisation

**Table/Fig-4B**: Clinical resemblance after a week of application of cryogen

**Table/Fig-5A**: Patient with pigmentation in the mandibular anterior region

**Table/Fig-5B**: Patient with pigmentation in the maxillary anterior region

**Table/Fig-6A**: Postoperative photograph after 6 months following gingival depigmentation

**Table/Fig-6B**: Postoperative photograph after 6 months following gingival depigmentation
absence of chronic inflammatory infiltrate in the connective tissue. Patients with pigmentation were treated using this technique and the results showed clinical applicability of this material [Table/Fig-4B]. Following 6 months of treatment there was clinically visible changes as compared to the results after depigmentation [Table/Fig-5A, 5B] and [Table/Fig-6A, 6B].

DISCUSSION

In the present study using ultralow temperature cryogen tetrafluoroethane on gingival biopsy showed that after 8 hours of application the epithelium showed degenerated rete pegs, and was elevated from the underlying connective tissue; this finding is similar to the finding in the study done by Mayer P D, Gerald Tussing, Frank M Wentz [11].

Studies have shown that superficial tissue necrosis occurs first after the cryosurgery then rapid epithelial migration covered the area [3]. Epithelial migration covers the denuded connective tissue and regenerate rapidly after a week of application. The stability of the depigmented area after six months of application of the cryogen is similar to the results seen in other studies [10, 12].

Gas expansion system used in cryosurgery is expensive and it is not widely used [10]. Cryosurgical treatment of oral lesions by liquid nitrogen [13,14] has been reported but both the gas expansion cryosurgical system and liquid nitrogen applied with cotton swabs are not easily obtainable in most clinics. Cross infection is also a disadvantage of using dipstick method [15].

In contrast to laser surgery and the conventional cryosurgery methods, the Tetrafluoroethane cryosurgery serves as an inexpensive, easy-to-use, store and transport [16]. In addition, the lack of bleeding and scar formation, its application without a regional anaesthesia, sutures or dressing, the ease of application of the cryogen at the papillary area by using a small cotton pledget, the added advantage of the depth control by using the time factor have added advantage of the depth control by using the time factor cryogen at the papillary area by using a small cotton pledget, the anaesthesia, sutures or dressing, the ease of application of the cryogen and the depth control by using the time factor 

CONCLUSION

The effect of ultralow temperature on the gingival tissue causes the epithelium to undergo cryonecrosis, helps to eliminate gingival melanin pigmentation by controlled epithelial destruction and clinically the gingiva heals without untoward side-effect and would be esthetically pleasing.

REFERENCES