COMPARATIVE EVALUATION OF EFFICACY OF GINGIVAL RETRACTION USING CHEMICAL AND MECHANICAL METHODS: AN IN VIVO STUDY

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Received: 19 September 2017, Revised and Accepted: 31 October 2017

ABSTRACT

Objective: The purpose of this in vivo study was to compare and evaluate the clinical efficacy of two gingival retraction systems; Ultrapak and Traxodent, based on the amount of gingival retraction achieved in vertical and horizontal direction and their hemorrhage control.

Methods: A total of 60 subjects were selected requiring fixed prosthesis. The two gingival retraction systems were used on the prepared abutments randomly. The vertical gingival retraction was measured before and after retraction using flexible measuring strip with 0.5 mm grading. The horizontal retraction was measured on the casts poured in polysilicone impressions made before the retraction and after retraction.

Results: Statistically significant difference (p<0.05) was found between the amount of the retraction (vertical and horizontal) achieved by Ultrapak as compared to Traxodent. However, in achieving hemostasis Traxodent showed better efficiency than Ultrapak (p<0.05).

Conclusion: The mean retraction width and depth achieved with retraction cord (Ultrapak) was significantly greater when compared with retraction paste. Although retraction paste (Traxodent) showed bleeding index significantly less when compared to that of retraction cord (Ultrapa).

Keywords: Gingival retraction, Ultrapak, Traxodent, Hemostatic control.

INTRODUCTION

Marginal adaptation plays an important role in the long-term success of the restoration, and failure to achieve the same can result in ill-fitting crowns, hypersensitivity, marginal leakage, periodontal tissue inflammation, and increased risk of recurrent caries [1,2]. The process of gingival displacement allows the exposure of the gingival or subgingival finish line along with the adjacent unprepared part of the tooth [3].

At present, in the market, there are various methods of the gingival tissue management such as mechanical method (retraction cords), chemo-mechanical method (chemicals embedded in cords), and surgical method (lasers, electrosurgery, and rotary curettage), of which gingival retraction cords are most commonly used [4,5].

The use of retraction cords can generate decent retraction, but clinicians usually report with the problem of gingival trauma and the time taken in the placement of the cord. Furthermore, there have been various investigations into the tendency of displacement cords to encourage bleeding and cause acute injury, which usually takes more than 1 week to heal. Gingival manipulation may result in significant bleeding in those patients taking antiplatelet medications [6-8] and those with pre-existing periodontal diseases [9]. Hence, the retraction material should not only displace the gingival tissue laterally and vertically but also control the bleeding [1,10-12].

Recently, cordless systems have been developed to save time and enhance patient compliance. The material offered is usually paste or foam that is injected into the crevicular sulcus [1,13,14]. This removes the need for the clinician to physically compress the material into the sulcus, where it may generate high pressure and cause injury.

The newly introduced Traxodent®hemodent paste retraction material (Premier Dental Products Co.) comprises 15% aluminum chloride topical paste and cotton caps, have been designed to improve the gingival displacement and assist in hemostasis.

Thus, the aim of the following study was to compare the efficiency of the gingival cord (mechanical) and Traxodent (chemical) in achieving the horizontal and vertical displacement of the gingival tissue along with its hemostatic potential.

METHODS

A total of 60 patients were selected for this study. Before the study, the protocol was explained to the participants, and informed consent was obtained. The study protocol was approved by the Institutional Ethical Committee. The sample size was further divided into two equal groups of 30 each, Group A and Group B.

1. Group A (30 patients): Gingival retraction cord Ultrapak 000 was used the mechanical method.
2. Group B (30 patients): Traxodent® retraction material was used the chemical method.

The inclusion criteria for the study included:
1. Any posterior (premolars and molars) edentulous area requiring a fixed partial denture.
2. Age >18 years.
3. Clinically and radiographically healthy abutment tooth.
4. Abutment teeth of normal size and contour (no development anomaly or regressive changes).
5. Probing depth <3 mm.

While the exclusion criteria were:
1. Age <18 years.
2. Signs of attachment loss and clinical mobility.
3. Presence of exudates.
4. Uncontrolled diabetes, hypertension, hyperthyroidism, and other cardiovascular disorders.

[Note: The tables and figures mentioned in the original text are not reproduced here due to the nature of the task to transcribe the natural text representation.]
Before starting the crown preparation, alginate impression was made for the selected arch to fabricate the customized sectional tray for making an elastomeric impression. This was followed by a veneer crown preparation where the finish line was placed equigingivally. After this, an elastomeric impression was taken with the help of addition polysilicon using the customized sectional tray using double mix single impression technique. The impression was checked for any voids, following which it was boxed and then poured in diestone. The cast retrieved was used to check for the horizontal retraction.

Then a smooth, flexible measuring strip with 0.5 mm grading was used to measure the sulcular depth in the vertical direction at mesiobuccal, midbuccal, and distobuccal region before the retraction was performed (Fig. 1 and 2). Next, the gingiva around the abutment tooth was retracted either with the help of Ultrapak (Group A) or Traxodent (Group B).

Retraction with the retraction cord (Ultrapack 000)

After measuring the sulcular depth, sufficient amount of retraction cord was cut and looped around the preparation. With the help of the cord packer, it was gently tucked down beneath the finish line of the preparation (Fig. 3). It was left in the place for 5 min.

Then, the cord was removed, and the preparation was air-dried. The hemostatic potential was checked, and accordingly, scores 0, 1, and 2 were given in Table 1.

Retraction with traxodent

The prepared tooth was air-dried, and it was made sure that the margins were kept dry. The paste was dispensed with the help of a syringe into the gingival sulcus. The paste was slowly applied into the sulcus all around the teeth (Fig. 4). The whitening of the gingiva showed that the paste was well applied. The paste was left there for 2 min (according to manufacturer's guidelines) and then rinsed off with water. Next, the tooth surface was air-dried, and hemostatic potential was checked, and accordingly, scores 0, 1, and 2 were given in Table 1.

After the retraction procedure (both in Group A and Group B) sulcular depth in the vertical direction using flexible strip (0.5 mm gradient) was measured in mesiobuccal, midbuccal, and distobuccal region, and post-retraction elastomeric impression was made, and the cast was retrieved.

The cast obtained (both pre- and post-retraction) was checked under the stereomicroscope (Fig. 5), and horizontal retraction was checked in mesiobuccal, midbuccal, and the distobuccal region (Fig. 6).

| Table 1: Hemorrhage scores |
|----------------------------|
| Score 0 | No bleeding |
| Score 1 | Bleeding controlled within 1 min |
| Score 2 | Bleeding not controlled within 1 min |

**Table 2: Mean values and standard deviations of horizontal and vertical retraction achieved in Group A and Group B (n=30)**

| Direction | Site       | Mean±SD   | p            |
|-----------|------------|-----------|--------------|
|           | Group A    | Group B   |              |
| Horizontal| Mesiobuccal| 0.24±0.038| 0.23±0.031   | 0.055; NS  |
|           | Midbuccal  | 0.26±0.038| 0.23±0.025   | <0.001; Sig|
|           | Distobuccal| 0.27±0.043| 0.23±0.023   | <0.001; Sig|
|           | Average    | 0.26±0.033| 0.23±0.019   | <0.001; Sig|
| Vertical  | Mesiobuccal| 1.01±0.334| 0.60±0.193   | <0.001; Sig|
|           | Midbuccal  | 1.00±0.331| 0.61±0.205   | <0.001; Sig|
|           | Distobuccal| 0.97±0.343| 0.55±0.157   | <0.001; Sig|
|           | Average    | 1.00±0.295| 0.59±0.157   | <0.001; Sig|

Sig: Significant, SD: Standard deviation, NS: Non significant

**Data analysis**

Statistical analysis was performed using SPSS version 18. p<0.05 was considered statistically significant. Comparison of mean scores for horizontal and vertical retraction was done using independent sample t-test, and Mann–Whitney U test was used to compare the hemostatic score.

**RESULTS**

**Fig. 1: Flexible measuring strip**

**Fig. 2: Measurement of the sulcular depth**

**Fig. 3: Placement of the cord**

**Fig. 4: Application of the Traxodent**
The mean horizontal and vertical retraction achieved by different retraction systems are listed in Table 2. The comparison of mean scores for horizontal and vertical retraction was done using independent sample t-test, and it showed a significant difference (p<0.05) in the amount of retraction achieved.

The hemorrhage scores on removal of each retraction system were compared using Mann–Whitney U test in Table 3 and the results showed a significant difference (p<0.05).

The Ultrapak retraction cord induced maximal bleeding on removal, while Traxodent induced minimal bleeding on removal.

DISCUSSION

Clinical parameters such as the location of the finish line, periodontal health of the patient along with sulcular bleeding during the process of impression directly influence the quality of the impression making in fixed partial prosthodontics. To obtain accurate marginal fit of the prosthesis a precise transfer from the patient to the definitive cast is essential. Hence, gingival displacement is necessary not only to record the gingival finish line but also the adjacent tooth structure [15].

CONCLUSION

This clinical study investigated the retraction efficiency and hemostatic potential of the cord and cordless system of gingival retraction. The mean retraction width and depth achieved with retraction cord (Ultrapak) was significantly greater when compared with retraction paste (Traxodent), while retraction paste showed bleeding index significantly less when compared to the retraction cord.

AUTHORS CONTRIBUTION

Parampreet Kohli-Compiled and analysed the data. Veena Hegde-Analyzed the data and prepared the manuscript.

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None.

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