Clinical study of a method for the receptor apparatus of the teeth protection at the stages of treatment with non-removable prosthesis designs

O. V. Voznyi*1,E,F, I. V. Yanishen2A,C, I. L. Diudina2A,B, V. H. Tomilin2B,E, A. V. Pohorila2D,E

1Zaporizhzhia State Medical University, Ukraine; 2Kharkiv National Medical University, Ukraine

Key words: receptor apparatus, intact teeth, electro-odontometry, mastication, protection technique, light-curing adhesive, antimototoxic drug.

This article presents the results of clinical approbation of our proposed method for protecting the receptor apparatus of the teeth through a complex of domestic light-curing adhesive preparations and antimototoxic drug “Traumeel”, the application of which was substantiated by experimental studies in laboratory animals.

The purpose of the study was to confirm clinically the experimentally obtained results to protect the receptor apparatus of the teeth by using the antimototoxic drug combined with the domestic light-curing adhesive.

Materials and methods. The method was tested in 72 patients, of whom 57 were the main group and 15 were controls. They were divided into 3 subgroups by age. A total of 200 teeth were prepared for one-piece cast fixed implant-supported prostheses. Electro-odontometry (EO) was performed and masticatory force (MF) was measured by the proposed method (patent number 99095142 dated September 16, 1999) before the preparation, when an anesthetic was worn off and one month after preparation.

Results. The positive results of our method were clinically confirmed by the EO data analysis and the MF values measured by proposed by us method on the day of the study, the next day, in a month after the intact teeth were coated with the proposed complex of preparations. When assessing the data obtained, it was found that there were no significant changes in the EO in indices as well as MF indicators on the day of the examination, the next day, in a month after the examination in the study group compared with the control.

Conclusions. Analyzing the results obtained, it can be noted that our method has a significant advantage in protecting the stumps during teeth preparation at the stages of treatment with non-removable prosthesis design and contributes to preventing complications of the hard dental tissue preparation.

Clinical испытание метода защиты рецепторного аппарата зубов на этапах лечения несъемными конструкциями протезов

О. В. Возний, И. В. Янишен, И. Л. Диудина, В. Г. Томилин, А. В. Погоріла

Мета роботи – клінічне підтвердження експериментально отриманих результатів щодо захисту рецепторів жувального тиску зубів шляхом застосування антимотоксичного препарату разом з вітчизняним світлотвердим адгезивом.

Матеріали та методи. Метод апробований на 72 пацієнтах, з них 57 утворили основну, а 15 – контрольну групу, яких поділили на 3 підрозділи з віковими ознаками. Препарували під опорні елементи незнімних суцільнолитих протезів 200 зубів. Вимірювали показники електроодонтометрії (ЕО) та жувального тиску (ЖТ), застосовуючи метод, який запропонували (патент № 99095142 від 16.09.1999), до операції препарування, після завершення дії аnestезії та через місяць після препарування.

Результати. Позитивні результати розробленого методу в клініці підтверджуються аналізом даних електроодонтометрії та значення жувального тиску, що вимірювали за методикою, яку запропонували, в день дослідження, наступного дня, через місяць і через рік після того, як інтахні зуби були покриті комплексом препаратів за допомогою методу, який запропоновали. Оцінюючи результати, встановили: в основній групі порівняно з контрольною показники електроодонтометрії в день дослідження, через день, через місяць після дослідження не мали значущих змін, те саме стосується показників жувального тиску в названих періодах дослідження.

Висновки. Аналізуючи отримані результати, визначили, що метод має суттєву перевагу щодо захисту кукси під час підготовки інтахніх зубів на етапах лікування незвідними конструкціями протезів і сприяє запобіганню ускладнень.

Клиническое испытание метода защиты рецепторного аппарата зубов на этапах лечения несъемными конструкциями протезов

А. В. Возный, И. В. Янишен, И. Л. Диудина, В. Г. Томилин, А. В. Погорелая

Цель работы – клиническое подтверждение экспериментально полученных результатов по защите рецепторов жевательного давления зубов путем применения антигомотоксического препарата с отечественным светоотверждающим адгезивом.
Materials and methods. The method was tested in 72 patients, 57 of whom were the main group and 15 were controls. They were divided into 3 subgroups by age. In total, 200 teeth were prepared for one-piece cast fixed implant-supported prostheses. The dental stump was prepared according to our method consisting of one-piece cast fixed implant-supported prostheses. The dental hard tissues were ground with a centered and sharp water-cooled abrasive tool at 300,000 revolutions per second. After preparation, the stumps of the teeth were covered with etching gel for 20–30 seconds. This significantly contributes to medicines penetration into the dental tissues. Then the gel was washed away by a water stream, sterile cotton pellet was placed to isolate the stump of the tooth and absorb any oral fluids and, if necessary, a saliva aspirator was used additionally, the stumps were dried with warm air. The antimicrobial drug “Traumeel” was applied evenly over the dental stump surface with an applicator and a current of warm air. The domestic light-curing adhesive was subsequently applied with the applicator over the antimicrobial preparation. The current of warm air was used to remove the rest of the drug. Following that, the adhesive was polymerized for 20 seconds. To reduce the environmental effect on the dental stumps of the prepared tooth, temporary crowns that were point-of-care made before the dental preparation, were fixed with the water dentine via material of GNJ Tempo Lux Company according to the standard methods [10, 11].

For clinical validation of the experimental study results, the parameters of EO and MF were measured according to the method proposed by us (patent No. 99095142 of 16.09.1999) before the preparation, when an anesthetic was worn off and one month after the preparation.

Results

The data analysis was aimed at determining the dynamics of EO and MF indicators, naturally occurring after the dental hard tissues preparation, the effectiveness of our proposed methodology in the experimental and control groups of patients as well as identifying the factors contributing to positive and, possibly, negative changes. ANOVA was used to measure statistical differences between results.

At present, the most commonly used prosthesis constructions are plastic- or ceramic-coated one-piece-cast, but the number of cases involving intact tooth pulp extraction, which is considerably supportive, regardless of an initial state of the tooth, has significantly increased in fixing these dentures. The reason for this is the desire of orthopedic dentists to avoid possible complications in the hard dental tissue preparation linked to the data about possible damage to the dentinal tubules and, as a result, the disturbance of hydrodynamic odontoblast processes. Depending on the removed layer of dentin, there is also a possibility to start compensatory mechanisms. However, it depends on the volume of tissues prepared, time interval since intervention [1,2,6,9,10,11,14].

Nevertheless, these protective reactions do not always start. Therefore, the first stage of prosthetic preparation is a dental pulp extraction. But some researches have proven that this manipulation significantly reduces the strength of dentin, which is the reason of frequent damage to the tooth crown after prosthetic treatment and a decrease in toughness and resistance to MF during functional loading [3,5,7,8,12].

To preserve intact teeth and prevent negative consequences, various methods of pulp extraction and techniques for preservation of supporting pulp vitality and reduction of their sensitivity after preparation were proposed by many scientists. According to these methods, the stumps of the prepared teeth were covered with various materials, provisional crowns, which were fixed with one-step and multi-step self-etching adhesives. However, none of the proposed methods resulted in restoration of the dental hard tissue and odontoblast processes damage during preparation, which are a part of the mechanoreceptors in the tooth pulp [4,13,15].

In view of the above and taking into account the relevance and practical importance of this issue, we proposed a method that was tested in the clinical practice of orthopedic dentistry.

Purpose

The purpose of the study was to confirm clinically the experimentally obtained results to protect the receptor apparatus of the teeth by using the complex of antimicrobial drug “Traumeel” and the domestic light-curing adhesive.
of the analysis using SAS software package. Regression analysis was implemented using Statistica software package (StatSoft Inc., № JPZ8041382130ARCN10-J).

As shown in Table 1, the EO parameters remained practically unchanged or slightly increased in the study group before the operation of preparation, on the next day and in a month after that when using our method, and did not depend on the group of teeth.

The EO values showed almost the same downward trend, except for incisors, after the preparation in both the study and control groups.

However, the picture was significantly changed in a month. If the EO indicators remained the same as they were before the preparation for the studied group, a statistically significant increase in these indices was observed in the control group concerning practically all anatomically oriented teeth, except for canines.

Looking at the average EO values in the study group, it can be seen that indicators were not increased in 84.96% of cases after the experimental processing of the teeth. In 13.54% of cases, there was a slight increase in this indicator and only in 1.5% of cases it was increased, that can be interpreted as a complication.

In the controls, unlike the study group, the EO index was not increased only in 20% of cases, and it was significantly increased in 23.3% of cases that can be interpreted as a complication, but a change in the indices was of 1–2 units from baseline in 56.7% of cases. It is notable that the control group indicators were more than 2 times higher than in the study group. This indicates a much less certainty regarding the control processing results.

There were only 2 cases with complications in the study group constituting 1.5% of the total number of teeth processed by our method. At the same time, there were 14 complicated cases in the control group accounting for 23.3% of the studied teeth number (Fig. 1).

The average values (M) of the MF and the standard errors of the mean (m), obtained from the clinical study, by patient groups and anatomically oriented groups of teeth are given in Table 2.

The analysis of MF measurements (Table 2) demonstrated independence of these indicators from the group of teeth and absence of their significant changes in the studied individuals.

The MF data from Table 2 shows that the increase in this index was statistically significant for not only the incisors

**Table 1.** Indicators of electro-odontometry, мA

| Group | Incisors | Canines | Premolars | Molars |
|-------|---------|---------|-----------|--------|
|       | Before | On the next day | In a month | Before | On the next day | In a month | Before | On the next day | In a month |
| Study | M      | 3.24    | 2.21      | 3.34    | 4.10    | 2.85      | 3.95    | 4.36    | 3.06      | 4.42    | 5.20    | 3.74      | 5.65    |
|       | m      | 0.09    | 0.09      | 0.15    | 0.23    | 0.21      | 0.25    | 0.15    | 0.12      | 0.14    | 0.16    | 0.18      | 0.24    |
|       | P1 < 0.001 | P2 > 0.05 | P1 < 0.001 | P2 > 0.05 | P1 < 0.001 | P2 > 0.05 | P1 < 0.001 | P2 > 0.05 | P1 < 0.001 | P2 > 0.05 | P1 < 0.001 | P2 > 0.05 |
| Control | M      | 3.22    | 2.67      | 6.22    | 4.75    | 3.50      | 5.25    | 4.20    | 2.93      | 6.07    | 4.83    | 3.89      | 6.89    |
|       | m      | 0.15    | 0.24      | 0.76    | 0.48    | 0.50      | 0.48    | 0.20    | 0.16      | 0.36    | 0.22    | 0.14      | 0.62    |
|       | P1 < 0.05 | P2 < 0.001 | P1 < 0.001 | P2 < 0.001 | P1 < 0.001 | P2 < 0.001 | P1 < 0.001 | P2 < 0.001 | P1 < 0.001 | P2 < 0.001 |

**P1:** significance between before the preparation and in a month after that.

**Table 2.** Masticatory force indicators, pF

| Group | Incisors | Canines | Premolars | Molar |
|-------|---------|---------|-----------|-------|
|       | Before | On the next day | In a month | Before | On the next day | In a month | Before | On the next day | In a month |
| Study | M      | 10.79   | 12.79     | 11.41  | 14.10    | 18.15     | 14.40   | 18.44   | 20.53     | 19.00    | 31.66   | 33.97     | 32.63   |
|       | m      | 0.23    | 0.26      | 0.40    | 0.46    | 0.47      | 0.47    | 0.39    | 0.39      | 0.38    | 0.59    | 0.57      | 0.81    |
|       | P < 0.001 | P > 0.05 | P < 0.001 | P > 0.05 | P < 0.001 | P > 0.05 | P < 0.001 | P > 0.05 | P < 0.001 | P > 0.05 | P < 0.001 | P > 0.05 |
| K     | M      | 10.22   | 12.55     | 13.44  | 15.00    | 17.55     | 16.75   | 18.89   | 21.83     | 22.00    | 33.00   | 35.67     | 36.27   |
|       | m      | 0.60    | 0.63      | 0.69    | 0.71    | 0.75      | 1.11    | 0.35    | 0.36      | 0.40    | 0.46    | 0.47      | 0.65    |
|       | P < 0.01 | P < 0.01 | P < 0.05 | P < 0.05 | P < 0.001 | P < 0.001 | P < 0.001 | P < 0.001 | P < 0.001 | P < 0.001 |

**P1:** significance between before the preparation and on the next day after that; **P2:** significance between before the preparation and in a month after that.
and canines on the next day after processing if compared between the study and control groups, as this increase concerning the premolars and molars was significantly greater in the control group than in the experimental group.

The MF indicators of the experimental group were not increased in 55.64 % of cases in a month after processing. In 42.1 % of cases, this indicator was increased of 1–2 pF and only in 2.26 % of cases – of 4 pF and more that can be understood as complications.

In the control group, the MF indicator was not increased only in 3.33 % of cases and its significant increase (4 pF and more) was observed in 36.66 % of cases that could be considered as complications (Fig. 2).

Discussion
From the analysis of the obtained results, it can be seen that the EO and MF indicators remained at the same level in a month after preparation in more than 50 % of the cases and did not depend on the tooth anatomical orientation in those patients who underwent the proposed by us protective method of stumps supporting during teeth preparation [12]. Nevertheless, in the control group, where our technique was not applied, the same indicators were significantly increased in a month and the anatomical dependence of the tooth was very important. The largest increase in these indicators was observed for incisors, then for canines, premolars and less for all the molars.

Conclusions
Thereby, we can see a significant advantage of our method in protecting the intact stumps during teeth preparation at the stages of treatment with non-removable prosthesis design. This makes it possible to apply our protective method of stumps supporting in the wide orthopedic practice and significantly increase the use of orthopedic non-removable denture helping prevention of complications in the hard dental tissues preparation.

Conflicts of interest: authors have no conflict of interest to declare.

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