A Randomized Split Mouth Clinical Trial Comparing Mineral Trioxide Aggregate with a New Fast-setting Calcium Silicate Cement in Direct Pulp Capping of Primary Molars: A Preliminary Report from a Long-term Follow-up

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ABSTRACT

Aims and objectives: This study was done to compare the success rate of a novel fast-setting calcium silicate cement (protooth) with mineral trioxide aggregate (MTA) in direct pulp capping (DPC) of primary molars.

Materials and methods: Forty-five patients with an age range of 5–7 years with 90 bilaterally symmetrical primary molars that had deep carious lesions were incorporated into a randomized split mouth clinical trial. Initially, the caries was removed. Afterward, the teeth randomly underwent DPC with either MTA or protooth. Restoration of the teeth was done by amalgam fillings. Clinical and radiographic examinations were performed after 6 months. To analyze the data, Chi-square statistical test was used. Values < 0.05 were considered statistically significant.

Results: After 6 months of follow-up, 88 teeth on 44 patients were available for evaluation. The MTA-treated teeth showed a success rate of 95.5%, while the same outcome for the protooth-treated teeth was 93.2%. The difference did not yield a statistically significant difference (p value > 0.05).

Conclusion: The findings of this study showed favorable results for the novel calcium silicate cement “protooth” when compared to the outcomes of MTA in the DPC of primary molars.

Clinical significance: The constant need for the development of more effective materials in the modern pediatric dentistry makes this novel cement of particular interest for pedodontists. The new cement is biocompatible, hydrophilic, and has fast-setting time and adequate tensile strength with favorable clinical results in the DPC of primary molars which can benefit pedodontists in their clinical practice.

Keywords: Calcium silicate cement, Direct pulp cap, Mineral trioxide aggregate, Primary molar, Protooth.

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INTRODUCTION

Preserving the vitality of primary teeth is one of the basic objectives in the practice of pediatric dentistry.1 This is because premature loss of deciduous teeth can lead to various malocclusions and esthetic problems.2,3 Conservative treatments like direct pulp capping (DPC) can considerably increase the durability of deciduous teeth in the mouth and minimize the need for more invasive and expensive treatments like pulpotomy or pulpectomy.4 Direct pulp capping includes the use of a biomaterial over the exposed pulp to preserve its vitality and optimally result in the formation of new dentine like bridges at the exposed site.5 However, DPC in deciduous teeth remains controversial because of its unsatisfactory success rate.6 The reason may be attributed to the higher amount of undifferentiated mesenchymal cells in the deciduous pulpal tissue which differentiate into odontoclasts in the presence of inflammation and bacteria, leading to internal resorption and the consequent failure of DPC.7 Therefore, selection of a material with favorable biocompatibility, antibacterial characteristics, and a tight seal which provokes dentinogenesis is crucial for a successful DPC treatment in the primary dentition.8

Thus far, various materials have been used in DPC technique in the literature. Calcium hydroxide has usually been the traditional first choice.9 However, the long-term solubility and gradual disintegration as well as the tunnel-like defects that are induced in the most portions of the tertiary dentinal bridges, compromise the long-term seal, and eventual success rate.10,11 Mineral trioxide aggregate (MTA), which is an alternative material, has been used in various pulp treatments including the DPC.12 It is a calcium silicate cement that tends to set in the humid oral environment.13 The calcium hydroxide, which releases as a by-product of hydration, results in some favorable properties like dentinogenesis in the pulpal tissue and apatite formation as well as antibacterial characteristics of the material.14

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Despite the successful use of MTA in the DPC of primary teeth, long setting time, difficult handling, and wash out of the material from the exposed site limits its clinical applications. This is especially important when encountering an exposed pulp that needs a fast-setting cement. In such situations, the drawback of MTA is emerged which requires a second appointment or having to cover up the material with an additional cement layer, e.g. glass ionomer to prevent wash out in one session treatment.

Recently, a new radiopaque and at the same time fast-setting and hydrophilic calcium silicate cement containing fluoride, named “protooth,” has been advocated to overcome the limitations of MTA. The mechanical properties of protooth is significantly superior to MTA and supports apatite formation in physiologic-like solutions. The setting time of protooth varies with respect to the consistency of the material (from creamy to thick condensable) and depending on its clinical application. The ultra-fast protooth which is intended for pulp capping purposes has a setting time of less than 2 minutes. The humid environment increases the cement’s mechanical characteristics and the biocompatibility of the material has been reported similar to the MTA.

Considering the favorable properties of this novel cement and the constant need for more effective material in the modern pediatric dentistry practice, this study was conducted to compare the efficacy of the “protooth” cement with MTA as the material of choice, both clinically and radiographically, in a split mouth clinical trial. The findings of this ongoing trial from which the preliminary findings are presented will help clarify the suitability of a new biomaterial in the practice of pediatric dentistry.

Materials and Methods

This study was a randomized split mouth clinical trial that was conducted at Tabriz University of medical sciences, between October 2017 and October 2018. The design of the study was similar to previous DPC trials. Before initiation of the study, the corresponding university research ethics committee reviewed and approved the protocol of the trial. The study was in compliance with the Helsinki Declaration of Human Rights. The current investigation was also listed at the Iranian Registry of clinical trials (IRCT ID: IRCT20100125003168N6). Patients and their parents were given adequate information about the possible risks and also the benefits of the study. Afterward, informed consents were obtained before the patients entering the trial.

Participants

Initially, dental examination was performed on a total of 200 patients referring to the department of pediatrics. From this initial examination, 45 patients (age range 5–7) with 90 bilateral deep carious second primary molar teeth in the same arch with the following inclusion criteria were incorporated into the study:

The patients had to be physically and mentally healthy, had no previous history of systemic disease, had no previous/current use of special systemic drugs, and had no allergic reactions. The bilateral primary molar teeth must have deep caries with vital pulp and should be restorable with amalgam. Furthermore, the teeth that had pathologic mobility, history of spontaneous pain, redness or swelling of vestibule, draining sinus tract, and sensitivity to palpation and/or percussion were removed from the trial. The teeth that had any sign of PDL widening, radiolucency on periapical or furcation area, or evidence of internal/external root resorption were also excluded from the study.

Sample Size

The sample size was estimated based on some previous DPC studies with MTA. Considering similar results with “protooth”, a power of 80%, and a two-sided significance level of 5%, 40 patients were estimated for the study. However, in order to increase the validity of the trial, and to compensate for fall outs from the research, 45 patients were listed in the study.

Clinical Procedure

In each participant, a carious second primary molar tooth was randomly allocated to the Protooth group (i.e., experimental group), while the counterpart tooth on the other side of the arch was allocated to the MTA group (control group). The authors flipped a coin for randomization of the teeth. One of the authors performed the procedures. Clinical and radiographic findings were registered both at the baseline and at the follow-up sessions. 2% lidocaine with 1/80,000 epinephrine (Daroupakhsh, Tehran, Iran) was used for achieving local anesthesia. A rubber dam was used for isolation purposes in each session. Diamond fissure bur (no. 008) on high-speed handpiece which was supplied by constant water irrigation was used for Enamel and peripheral caries removal. Dentinal caries was also removed using tungsten carbide round bur on low-speed handpiece. To wash away dentinal debris, the cavity was constantly irrigated using NaOCL 1%. In cases where the exposed pulps were pinpoint (i.e., <1 mm), surrounded by sound healthy dentin and the bleeding arrested in 2–3 minutes, DPC was performed. After pinpoint exposure, 0.12% chlorhexidine was used for washing the cavity and to reduce the bacterial load. Hemostasis was then achieved using a cotton pellet moisturized with sterile normal saline and a gentle pressure. A dry cotton pellet was then used to dry the cavity. If the bleeding did not stop in 2–3 minutes, or the exposure site was beyond 1 mm, the tooth underwent more aggressive treatments (i.e., pulpotomy or pulpectomy) and was excluded from the study. In cases in which pulpal tissues were not exposed after caries removal, conventional amalgam restoration was done, and the teeth were excluded from the trial.

In each tooth, after hemostasis control, the teeth randomly underwent treatment with either protooth or MTA according to the instructions of the manufacturer. Because of the different manipulation techniques for each study group, the operator was not blind to the material used. All other contributors were blinded to the study. A separate session was allocated for the treatment of the counterpart tooth in order to avoid discomfort caused by bilateral anesthesia. A round ended instrument (S Ball Burnisher 27/27S, Premier, USA) was used for application of Protooth (Dentsolve, Aarhus, Denmark) in 2 mm thickness over the exposed site. The material was extended approximately 1 mm peripherally beyond the exposed site to ensure complete covering. The material was further covered by a wet cotton pellet for 3 minutes to allow the setting process to occur. MTA carrier (G Hartzell & Son, #ISS52, 1.8 mm) was used to apply the MTA in thickness of 2 mm. The material was also extended approximately 1 mm beyond the exposed site for the same covering purposes. A layer of low-viscosity glass ionomer (Fuji II, GC, Tokyo, Japan) with 2 mm thickness covered the MTA to ensure the seal of the material. In both the groups, the teeth were alternatively restored using amalgam build up (Cinalux, Tehran Iran). A single session was used to perform all the procedures. Immediately after treatment, periapical radiographs were obtained. The radiographs were also repeated after 6 months of follow-up.
The flow diagram of the study from baseline up to the end of the follow-up period is described at Flowchart 1.

At the end of the follow-up, the teeth underwent clinical and radiographic examination by two expert pedodontists who were blinded to the material and were calibrated in a separate session before the study. All the radiographs were obtained by parallel technique. The presence of any of the following findings deemed failure of the treatment: Sensitivity to percussion, internal/external root resorption, widening of periodontal ligament, spontaneous pain, sinus tract, swelling, and intraradicular radiolucency or periapical lesions. In cases of disagreement, a third pedodontist was asked to perform the examination. The interexaminer reliability was verified using kappa agreement coefficient. In cases of treatment failure, the teeth underwent a more advanced pulpal treatment (i.e., pulpectomy).

Statistical Analysis
The descriptive data were tabulated for statistical analysis. In order to compare the treatment outcomes between different study groups, Chi-square analysis was utilized. The statistical analyzes were performed by SPSS 18 under windows. The statistical significance was set at $p$ values < 0.05.

Results
Forty-five patients (22 boys, 23 girls, mean age 7–8 years) were enrolled into the study. The interexaminer agreement at the end of the follow-up was excellent. ($\kappa = 0.92$, $p$ values < 0.001). After 6 months, a total of five teeth failed with two of the failed cases being in the MTA-treated group (4.5%) and three teeth in the protooth-treated group (6.8%). In addition, one patient dropped out of the study due to relocation to other city and therefore was excluded from the research. The failed primary molars at both MTA and protooth group underwent further complimentary treatment of pulpectomy with amalgam restoration. The failure reasons for both the MTA and the protooth-treated teeth are described in Table 1. Sensitivity to percussion demonstrated the highest reason of failure. None of the failed cases showed root resorption as the reason of failure. Chi-square statistical analysis yielded no significant difference between the success rate of MTA-treated teeth (95.5%) with that of protooth-treated cases (93.2%) at the end of the follow-up session (Fig. 1, $p$ values > 0.05).

Discussion
Direct pulp capping in primary teeth has always been a challenging procedure in pediatric dentistry. Because of its lower success rate in some clinical situations, some authors have questioned the procedure.24 The principal biologic difference between primary and permanent teeth is that the primary teeth have a highly populated undifferentiated mesenchymal cells. In the presence of inflammation caused by bacterial invasion, this potential can induce mesenchymal cells to differentiate into odontoclasts which subsequently can lead to internal root resorption.7 Therefore, it is essential to follow strictly the indications and contraindications of the DPC. A good case selection and strict adherence to antibacterial load reduction is of utmost importance in any successful DPC treatment. In this trial, the infected dentine was thoroughly removed. The case selection procedure was meticulously adhered and was strictly limited to those of less than 1 mm exposure with hemostasis control under 3 minutes. Furthermore, constant hypochlorite irrigation was used to reduce the microbial load before exposure and 0.12% chlorhexidine rinse after pulpal exposure to further augment the antiseptic procedure. These considerations might have contributed to the fact that no evidence of root resorption was discovered in any of the treated cases in our study. This was comparable to some recent clinical trials of DPC in the primary molars.4,21,22 However, while these principals are prerequisite to

| Evaluation criteria | MTA group, $n = 44$ | Protooth group, $n = 44$ |
|---------------------|--------------------|------------------------|
| Spontaneous pain    | 1                  | 1                      |
| Tenderness to       | 1                  | 2                      |
| percussion          |                    |                        |
| Sinus tract         | 0                  | 0                      |
| Root resorption     | 0                  | 0                      |
| PDL widening        | 0                  | 1                      |
| Overall failure rate| 2 (4.5%)           | 3 (6.8%)               |

Fig. 1: Success rates of mineral trioxide aggregate (MTA) and protooth treated teeth
successful DPC, other characteristics are also necessary for the capping material. The material should be biocompatible and have good physical, antimicrobial, and sealing properties. Calcium hydroxide has been the traditional pulp capping material. However, recently calcium silicate cements like MTA have gained increasing popularity with better long-term outcomes. Recent studies have shown the superior properties of MTA in comparison to calcium hydroxide in terms of sealing ability, physical properties, and biocompatibility. However, despite acceptable clinical performance of MTA, some drawbacks have also been reported for the material. Poor handling, long setting time, and wash out of the material are among the mentioned impediments. Because of these considerations, we decided to select and compare MTA as the control group with a new calcium silicate cement named “protooth”. The Protooth is a newly introduced fast-setting cement that contains fluoride. It has several potential applications in tooth crowns. The cement’s composition is similar to that of MTA, which includes calcium sulfate, tricalcium silicate, tricalcium aluminate, and dicalcium silicate. Additionally, the new cement contains radiocontrast material, fluoride additive, and nanosilica.

Appropriate mechanical properties and durability of a cement in humid oral environment are crucial in its ultimate clinical success. Therefore, the physical properties of protooth has been investigated in a previous laboratory study. The study showed a significantly higher early tensile strength as an indicator of mechanical properties of protooth when compared to the MTA or Biodentine. The ultimate strength of MTA did not show a significant difference than protooth, although a non-significant increase was observed in favor of protooth. Furthermore, it was shown that the mechanical properties of protooth cement significantly increases in humid environment over the course of time, a feature that is favorable for a pulp capping material. This hydrophilic characteristic is of particular interest for the pediatric purposes, since humidity control and isolation are challenging in such patients.

The apatite formation ability has been known to be an important indicator for biocompatibility of dental materials. This is because superficial formation of apatite layer may yield a suitable surface for cells to differentiate into odontoblasts. Apatite layer can also enhance the sealing ability of dentinal tubules as well as increase the material bonding to the dentine. In a previous laboratory study to investigate the biocompatibility of the protooth cement, the material was allocated into three compositions, protooth, ultrafast protooth, and high fluoride protooth. The principal difference between these compositions was the fluoride and the radiocontrast concentrations. The study showed that all types of the protooth have the ability to form apatite layer and that the thickness of the layer increases over the course of time resulting in more voluminous and compact morphological structures. Furthermore, it was shown that the “high fluoride” protooth can form a thicker apatite layer, since fluoride concentrations can increase the appetite-forming ability of the cement. However, because a fast-setting cement for the exposed vital pulp of primary teeth is essential, and because of cooperation issues of pediatric patients, the ultrafast protooth was selected for this study. This type of protooth can set in less than 2 minutes which is in clear advantage compared to the MTA. The mechanism of the excellent biocompatibility of the cement, is attributed to a hydration by-product of the cement, namely, calcium hydroxide, which reacts with the available phosphate ions of physiologic solutions resulting in the formation of calcium phosphates including apatite. It is suggested that the protooth-induced calcified bridges undergo a histological evaluation and comparison in future laboratory studies so that the quality of the induced appetite layer can be compared to that of MTA or calcium hydroxide materials.

The release of calcium hydroxide is known to render antibacterial properties to the material. Cytotoxicity of the biomaterials contacting directly the exposed pulp is an important factor when dealing with vital pulp therapy procedures like the DPC. In a previous invitro study on mouse fibroblasts, the cytotoxicity of several biomaterials were evaluated. The findings indicated that calcium silicate cements like “protooth” had a significantly lower cytotoxicity than calcium hydroxide based materials. Furthermore, there was no significant difference between the cytotoxicity of various calcium silicate based cements like the protooth, MTA, and Biodentine. Interestingly, the different compositions of protooth had almost an identical cytotoxicity level when compared with each other. In a recent case report, the authors reported acceptable tolerance and biocompatibility by human periapical tissues when using protooth as root end filling material in an avulsed open apex permanent incisor.

Despite existence of a number of in vitro studies on biocompatibility and mechanical properties of protooth, to our best knowledge, this is the first clinical trial evaluating this novel cement for vital pulp therapy procedures like the DPC. The findings from the current trial, showed favorable initial results with protooth when used in direct pulp capping of deciduous molars indicating that the difference between success rate of Protooth cement and the MTA was not statistically significant. However, these findings should be interpreted with caution since the current data are preliminary and longer follow-up times seem relevant in order to derive a more definitive conclusion. It is noteworthy that the current study will continue for long-term and also for other indications so that more information can be gathered from this novel biomaterial.

The successful use of MTA in DPC of deciduous molars is well documented. In our study, we found comparable clinical outcomes for this novel cement when compared to the MTA in the DPC of primary molars at short-term level. The new calcium silicate cement also had some further advantages to the MTA, including fast-setting time, better physical properties, and easy handling of the material. The findings of our study are in harmony with some recent clinical trials on the DPC of primary molars. Despite the highly populated mesenchymal cell content of primary molars, the DPC can be a successful conservative treatment if proper case selection is followed, a favorable pulp material is used and the correct procedure is meticulously adhered. However, further clinical trials with longer follow-up periods and larger samples sizes, along with more histological investigations into the material characteristics of this novel cement, seem relevant to derive more definite conclusions on the successful use of protooth cement for the DPC of primary molars.

**Conclusion**

Within limitation of the current study, it can be concluded that “protoooth” can successfully be applied in the DPC of deciduous molars. A proper case selection and meticulous adherence to the guidelines of DPC are recommended.
Novel Calcium Silicate Cement in the DPC of Primary Molars

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