

Clinical Trial Protocol

Iranian Registry of Clinical Trials

16 Sep 2026

Comparison clinical and radiographic success rate of TheraCal-LC® and Mineral Trioxide Aggregate in pulpotomy of primary teeth with 12 month follow up: A split-mouth randomized controlled clinical Trial

Protocol summary

Study aim

The effect of TheraCal-LC® and Mineral Trioxide Aggregate in pulpotomy of primary molars will be comprised with the aim of preservation of tooth vitality and structure and prevention of more aggressive treatments.

Design

In this randomized, single blind, split mouth, controlled clinical trial study with parallel group, symmetrical bilateral primary molar teeth in maxilla or mandible will be chosen in 45 patients (overall 90 teeth). Randomization will be done by throwing a coin.

Settings and conduct

This clinical trial study will be done in Tabriz dentistry faculty. After obtaining informed consent from patient's parents, local anesthesia injection, isolation, and caries excavation, in group A, TheraCal-LC® placement and in group B, MTA placement in exposure site will be performed. In this single blind study, participants are unaware of the allocation of study groups and information is not given to the parents about which of the materials is used in the patient's left or right primary molar.

Participants/Inclusion and exclusion criteria

Inclusion criteria: 5 to 8 years old children having a pair of symmetric primary molar teeth in one jaw having deep caries and vital pulp and no history of spontaneous pain; No sign of radiolucency or pathologic root resorption
Exclusion criteria: Lack of informed consent by the child patient's parent; Abnormal bleeding at the site of pulp exposure.

Intervention groups

Local anesthesia injection, isolation, and caries excavation and access cavity will be performed in both groups and in group A, TheraCal-LC® placement and in group B, MTA placement in exposure site will be performed. Then in both groups the site will be covered

with Glass Ionomer. After amalgam restoration, periapical radiographs will be obtained.

Main outcome variables

Clinical success (Dental pulp vitality after treatment);
Radiographic success.

General information

Reason for update

Acronym

است. نام SSC سه حرف Stainless Steel Crowns نام اختصاری
استو MTA سه حرف Mineral Trioxide Aggregate اختصاری

IRCT registration information

IRCT registration number: **IRCT20100125003168N7**

Registration date: **2020-03-03, 1398/12/13**

Registration timing: **prospective**

Last update: **2020-03-03, 1398/12/13**

Update count: **0**

Registration date

2020-03-03, 1398/12/13

Registrant information

Name

Leila Erfanparast

Name of organization / entity

Tabriz University of Medical sciences

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Recruitment status

Recruitment complete

Funding source

Expected recruitment start date

2020-04-05, 1399/01/17
Expected recruitment end date
 2020-04-15, 1399/01/27
Actual recruitment start date
 empty
Actual recruitment end date
 empty
Trial completion date
 empty

Scientific title
 Comparison clinical and radiographic success rate of TheraCal-LC® and Mineral Trioxide Aggregate in pulpotomy of primary teeth with 12 month follow up: A split-mouth randomized controlled clinical Trial

Public title
 Comparison success rate of TheraCal-LC® و Mineral Trioxide Aggregate in pulpotomy of primary teeth

Purpose
 Treatment

Inclusion/Exclusion criteria
Inclusion criteria:
 -Complete physical and mental health, with no confounding history of allergic reactions and systemic disease and/or use of special local or systemic drugs
 Having a pair of symmetric primary molar teeth (first or second) in one jaw, having deep caries and vital pulp which is capable of being restored by SSC
 No history of spontaneous pain, pathologic mobility, draining sinus tract, redness or swelling of vestibule
 Normal gingival and periodontal condition, with no sensitivity to vestibular palpation, and no pain on percussion test
 5-8 years old children
 No sign of radiolucency in periapical or furcation area
 No widening of PDL space or loss of lamina dura continuity
 No evidence of internal/external pathologic root resorption
Exclusion criteria:
 Lack of informed consent by the child patient's parent-
 Abnormal bleeding at the site of pulp exposure (i.e. bleeding with dark color which lasts for more than 5 minutes despite application of wet cotton pellet with mild pressure)

Age
 From 5 years old to 8 years old

Gender
 Both

Phase
 N/A

Groups that have been masked

- Participant

Sample size
 Target sample size: 45
 More than 1 sample in each individual
 Number of samples in each individual: 2
 90 teeth in 45 patients

Randomization (investigator's opinion)
 Randomized

Randomization description
 Randomization is done by throwing a coin. One side of coin is assigned to the MTA and the other side is

assigned to TheraCal-LC® and the first throw, determines the material used on the right primary molar of the patient.

Blinding (investigator's opinion)
 Single blinded

Blinding description
 In this study, participants are unaware of the allocation of study groups and information is not given to the parents about which of the materials is used in the patient's left or right primary molar.

Placebo
 Not used

Assignment
 Parallel

Other design features

Secondary Ids

empty

Ethics committees

1

Ethics committee

Name of ethics committee

Ethics Committee of Research Department - Tabriz University of Medical Sciences

Street address

Research Department- Tabriz University of Medical Sciences- Golgasht street- Tabriz- Iran

City

Tabriz

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East Azarbaijan

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5166614711

Approval date

2020-02-29, 1398/12/10

Ethics committee reference number

IR.TBZMED.RES.1398.1252

Health conditions studied

1

Description of health condition studied

Reversible Pulpitis

ICD-10 code

K04.0

ICD-10 code description

Pulpitis

Primary outcomes

1

Description

Dental pulp vitality after treatment

Timepoint

6 & 12 months

Method of measurement

Lack of any of clinical or radiographic signs or symptoms of treatment failure

dent.fac@tbzmed.ac.ir

Secondary outcomes

1

Description

Radiographic success

Timepoint

6&12 months

Method of measurement

Presence of one or more of the following radiographic signs will be considered as failure of treatment: internal and/or external root resorption, periodontal space widening, inter-radicular radiolucency, and periapical lesions.

2

Description

Clinical success

Timepoint

6 &12 months

Method of measurement

Presence of one or more of the following clinical signs/symptoms will be considered as failure of treatment: spontaneous pain, swelling, pathologic mobility, tenderness to pressure, and presence of sinus tract, swelling, and sensitivity to percussion.

Intervention groups

1

Description

Intervention group:

Category

Treatment - Other

Recruitment centers

1

Recruitment center

Name of recruitment center

Department Of Pediatric Dentistry, Tabriz University of Medical Sciences

Full name of responsible person

Dr Leila Erfanparast

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Sponsors / Funding sources

1

Sponsor

Name of organization / entity

Tabriz University of Medical Sciences

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Grant name

Grant code / Reference number

Is the source of funding the same sponsor organization/entity?

Yes

Title of funding source

Tabriz University of Medical Sciences

Proportion provided by this source

100

Public or private sector

Public

Domestic or foreign origin

Domestic

Category of foreign source of funding

empty

Country of origin

Type of organization providing the funding

Academic

Person responsible for general inquiries

Contact

Name of organization / entity

Tabriz University of Medical Sciences

Full name of responsible person

Dr Leila Erfanparast

Position

Associate Proffessor of Pediatric Dentistry department

Latest degree

Specialist

Other areas of specialty/work

Dentistry

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Sharing plan

Deidentified Individual Participant Data Set (IPD)

Yes - There is a plan to make this available

Study Protocol

Yes - There is a plan to make this available

Statistical Analysis Plan

Yes - There is a plan to make this available

Informed Consent Form

Yes - There is a plan to make this available

Clinical Study Report

Yes - There is a plan to make this available

Analytic Code

Yes - There is a plan to make this available

Data Dictionary

Yes - There is a plan to make this available

Title and more details about the data/document

The whole data is shareable.

When the data will become available and for how long

Access granted 6 months after publication of results

To whom data/document is available

Physicians and researchers in medical sciences as well as those involved in industry

Under which criteria data/document could be used

Data are released for non commercial purposes and for research objectives only.

From where data/document is obtainable

Dr.Sedigheh Hasanpour, Pediatric dentistry department, Tabriz dental faculty, Tabriz University of medical sciences, Golgasht street. s_hasanpour87@yahoo.com 04133346977

What processes are involved for a request to access data/document

The applicant should forward their application to the email provided. the data will be send via email at 72 hours at most.

Comments