

# Clinical Trial Protocol

## Iranian Registry of Clinical Trials

29 Jun 2026

### Comparison between acellular dermal matrix and connective tissue graft on vertical soft tissue thickness during submerged implant insertion

#### Protocol summary

##### Study aim

determination of alloderm and Connective Tissue Graft (CTG) on vertical soft tissue thickness on two-stage implant insertion

##### Design

Randomized two-arm parallel, double-blind trial, sample size 20 (10 patients in each group)

##### Settings and conduct

In this study, 1 hour before surgery, the patient receives 1 gram of prophylactic dose of amoxicillin, the procedure is then continued at 500 mg 3 times a day for 1 week after the intervention. The entire surgical procedure is performed by a surgeon. Buccal flap elevates with full thickness flap and vertical soft tissue dimension measured. Perio probe insert on alveolar ridge on implant insertion site. If vertical dimension of soft tissue was more than 3mm the patient went out of the study. After implant insertion in first group connective tissue graft (CTG) placed and in the second group ADM was placed in 10\*10 mm dimension on alveolar ridge due to randomization. Flap sutured with 0/4 silk suture. And after 3 month recalling the patient happened and on healing abutment placement measurement of soft tissue thickness evaluated and reported by the blinded examiner who was not aware of belonging of patient to each groups.

##### Participants/Inclusion and exclusion criteria

1-patients candidate for implant treatment 2-thin crestal mucosal tissue (2 mm or less); partial edentulousness in different areas of the mouth 3-no bone augmentation process before and during implant placement.

##### Intervention groups

CTG and alloderm insertion on a two-stage implant for evaluation of these materials on progression and improvement of soft tissue around dental implants

##### Main outcome variables

soft tissue thickness around implant

#### General information

##### Reason for update

##### Acronym

##### IRCT registration information

IRCT registration number: **IRCT20211003052658N1**

Registration date: **2021-11-07, 1400/08/16**

Registration timing: **retrospective**

Last update: **2021-11-07, 1400/08/16**

Update count: **0**

##### Registration date

2021-11-07, 1400/08/16

##### Registrant information

##### Name

amin mohammadpourkolde

##### Name of organization / entity

##### Country

Iran (Islamic Republic of)

##### Phone

+98 13 4212 5497

##### Email address

aminmpkld71@gmail.com

##### Recruitment status

**Recruitment complete**

##### Funding source

##### Expected recruitment start date

2021-05-23, 1400/03/02

##### Expected recruitment end date

2021-08-29, 1400/06/07

##### Actual recruitment start date

2021-07-24, 1400/05/02

##### Actual recruitment end date

2021-10-19, 1400/07/27

##### Trial completion date

2021-10-23, 1400/08/01

##### Scientific title

Comparison between acellular dermal matrix and connective tissue graft on vertical soft tissue thickness during submerged implant insertion

#### Public title

Comparison between acellular dermal matrix and connective tissue graft on vertical soft tissue thickness during submerged implant insertion

#### Purpose

Treatment

#### Inclusion/Exclusion criteria

##### Inclusion criteria:

Thin crestal mucosal tissue (2 mm or less) Partial edentulousness in different areas of the mouth No bone augmentation process before and during implant placement

##### Exclusion criteria:

Patients undergoing soft and hard tissue regeneration treatments Implant loss in the area Patients under 18 years of age Has a medical contraindication for implants Has a history of periodontitis, has a history of uncontrolled diabetes and smoking and alcohol consumption Systemic disease that interferes with implant placement, history of radiation therapy and chemotherapy

#### Age

From **18 years** old

#### Gender

Both

#### Phase

N/A

#### Groups that have been masked

- Participant
- Outcome assessor

#### Sample size

Target sample size: **20**

Actual sample size reached: **20**

#### Randomization (investigator's opinion)

Randomized

#### Randomization description

Individuals who have been included in the study based on inclusion and exclusion criteria, randomly allocate into control and test group through randomized block method. Each block will contain 2,4 or 8 patient. After completion of each block treatment will be done by a randomized permutation on patient. Blocks and permutations on each block will create by agricolae package in R software with a definite seed. The number of the seed provide the possibility of recreation of random sequence. regard to this package number of block, size of block, sequence in each block and type of each patient treatment will present as outcome of the software.

#### Blinding (investigator's opinion)

Double blinded

#### Blinding description

The patients included in the study did not know that they belonged to the control and intervention groups. And measurements are made by another person during the first and second stages who does not know that the treated people belong to the control and intervention

groups.

#### Placebo

Not used

#### Assignment

Parallel

#### Other design features

## Secondary Ids

empty

## Ethics committees

### 1

#### Ethics committee

##### Name of ethics committee

Ethics committee of Qazvin University of Medical Sciences

##### Street address

Qavin Shahid bahonar bolvar .Qazvin univercity of medical science

##### City

Qazvin

##### Province

Qazvin

##### Postal code

3419915315

#### Approval date

2021-07-20, 1400/04/29

#### Ethics committee reference number

ir.qums.rec.1400.187

## Health conditions studied

### 1

#### Description of health condition studied

mucosal thinness in relation to peri-implant disease

#### ICD-10 code

#### ICD-10 code description

## Primary outcomes

### 1

#### Description

soft tissue thickness

#### Timepoint

on implant insertion and on healing abutment placement

#### Method of measurement

with periodontal probe and gage guide

## Secondary outcomes

empty

## Intervention groups

### 1

#### Description

Intervention group 1: CTG placement  
**Category**  
Treatment - Other

2

**Description**  
Intervention group 2: ADM placement  
**Category**  
Treatment - Other

## Recruitment centers

1

**Recruitment center**  
**Name of recruitment center**  
Qazvin university of medical science  
**Full name of responsible person**  
Amin Mohammadpour Kolde  
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## Sponsors / Funding sources

1

**Sponsor**  
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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**  
Qazvin 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**  
Qazvin University of Medical Sciences  
**Full name of responsible person**  
Amin Mohammadpour Kolde  
**Position**  
Resident  
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## Person responsible for scientific inquiries

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## Person responsible for updating data

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**Province**

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## Sharing plan

**Deidentified Individual Participant Data Set (IPD)**

Undecided - It is not yet known if there will be a plan to make this available

**Study Protocol**

Undecided - It is not yet known if there will be a plan to make this available

**Statistical Analysis Plan**

Undecided - It is not yet known if there will be a plan to make this available

**Informed Consent Form**

Undecided - It is not yet known if there will be a plan to make this available

**Clinical Study Report**

Not applicable

**Analytic Code**

Not applicable

**Data Dictionary**

Not applicable