

# Clinical Trial Protocol

## Iranian Registry of Clinical Trials

10 Jun 2026

### Clinical trial comparing the effect of conventional method and method of use three-dimensional printers on the reconstruction difficulties of the mandibular body during surgery and afterwards

#### Protocol summary

##### Summary

20 adult patients, regardless of sex and age limits, from the patients referred to the department of oral and maxillofacial surgery of Yazd are selected. All of them must have benign pathological lesion (ameloblastoma, odontogenic keratocyst, etc) involving only mandibular body (from distal aspect of canine tooth to mandibular angle) unilaterally, requiring resection and subsequent reconstruction of the body of mandible under general anesthesia. Patients are randomly divided into two groups. They will be ready for operation, one group using conventional methods and the other using three-dimensional printing technology. In the group using three-dimensional printer, after acquiring a CT scan of axial, coronal and three-dimensional reconstruction views, prototype three-dimensional models of the patient's facial bones are fabricated. Then, while studying the three-dimensional model, required treatment plan is determined and shaping and adapting reconstruction plates on the model, which represents the state of the mandibular bone lesion morphology is done. The pre-shaped and adapted reconstruction plates will be delivered to hospital's CSR for sterilization. In the group using conventional method, CT Scan of axial, coronal and three-dimensional reconstruction views are provided for patients but required treatment plan is determined using only clinical assessment and radiographic evaluation and three-dimensional models will not be manufactured. After routine tests surgery is performed. The amount of mouth opening will be measured in both groups before surgery. In the intervention group, during surgery, after resection of bone lesions, pre-shaped plates are placed at the site of reconstruction; while in the control group reconstruction plates are shaped routinely during the surgery and adapted on the bone. Finally, after placement and fixation of reconstruction plates, surgical site will be

sutured to end the surgery. During surgery, the amount of blood loss and duration of the operation is measured in both groups. After surgery, mouth opening and post-operative pain is evaluated in multiple time periods. Also the risk of plate exposure and infection during six months of follow-up will be examined.

#### General information

##### Acronym

##### IRCT registration information

IRCT registration number: **IRCT201605073398N4**

Registration date: **2016-05-20, 1395/02/31**

Registration timing: **retrospective**

Last update:

Update count: **0**

##### Registration date

2016-05-20, 1395/02/31

##### Registrant information

##### Name

Alireza Navabazam

##### Name of organization / entity

Yazd University of Medical Sciences

##### Country

Iran (Islamic Republic of)

##### Phone

+98 353624760

##### Email address

navabazam@ssu.ac.ir

##### Recruitment status

**Recruitment complete**

##### Funding source

Shahid Sadoghi University Of Medical Sciences

##### Expected recruitment start date

2015-07-23, 1394/05/01

**Expected recruitment end date**

2015-09-23, 1394/07/01

**Actual recruitment start date**

empty

**Actual recruitment end date**

empty

**Trial completion date**

empty

**Scientific title**

Clinical trial comparing the effect of conventional method and method of use three-dimensional printers on the reconstruction difficulties of the mandibular body during surgery and afterwards

**Public title**

Compare the conventional method and the method of using three-dimensional printers in reconstruction of the mandibular body and Determine the effect of these methods on Intraoperative and postoperative problems

**Purpose**

Treatment

**Inclusion/Exclusion criteria**

Inclusion criteria: They all have a benign pathological lesion that only involve unilateral body of the mandible (the distal of canine tooth to the angle of mandible) and require resection and subsequent reconstruction surgical procedure under general anesthesia Exclusion criteria: Systemic problems; kidney defects; neurological and psychological problems; diabetes; hypertension; pregnancy; coagulation problems; use of bisphosphonates; the broad-spectrum antibiotics; corticosteroids and non-steroidal analgesics

**Age**

From **20 years** old to **70 years** old

**Gender**

Both

**Phase**

2

**Groups that have been masked**

*No information*

**Sample size**

Target sample size: **20**

**Randomization (investigator's opinion)**

Randomized

**Randomization description****Blinding (investigator's opinion)**

Not blinded

**Blinding description****Placebo**

Not used

**Assignment**

Parallel

**Other design features****Secondary Ids**

empty

**Ethics committees****1****Ethics committee****Name of ethics committee**

Ethics committee of Shahid Sadoghi University of Medical Sciences

**Street address**

Shahid Sadoghi University of Medical Sciences, Western Sadoughi Blvd, Bahonar Square

**City**

Yazd

**Postal code****Approval date**

2016-05-09, 1395/02/20

**Ethics committee reference number**

IR.SSU.REC.1395.20

**Health conditions studied****1****Description of health condition studied**

pathologic lesion of mandibular body

**ICD-10 code**

D 16.5

**ICD-10 code description**

Benign neoplasm of bone and articular cartilage

**Primary outcomes****1****Description**

surgery duration

**Timepoint**

from preparation of patient until extubation

**Method of measurement**

Questionnaire

**2****Description**

pain

**Timepoint**

2,6,12,24,48,72 hours after surgery

**Method of measurement**

Questionnaire

**3****Description**

duration of resection of bone and reconstruction of mandible

**Timepoint**

The onset of bone resection to close the last screw on the reconstruction plate

**Method of measurement**

Questionnaire

**4****Description**

blood loss

**Timepoint**

Intraoperative

**Method of measurement**

Questionnaire

**5****Description**

mouth opening

**Timepoint**

before surgery,24,48,72 and a week after surgery

**Method of measurement**

Questionnaire

**Secondary outcomes**

empty

**Intervention groups****1****Description**

intervention1: In the group using three-dimensional printer, after acquiring a CT scan of axial, coronal and three-dimensional reconstruction views, prototype three-dimensional models of the patient's facial bones are fabricated. Then, while studying the three-dimensional model, required treatment plan is determined and shaping and adapting reconstruction plates on the model, which represents the state of the mandibular bone lesion morphology is done. The pre-shaped and adapted reconstruction plates will be delivered to hospital's CSR for sterilization. After routine tests surgery is performed. The amount of mouth opening will be measured before surgery. during surgery, after resection of bone lesions, pre-shaped plates are placed at the site of reconstruction; Finally, after placement and fixation of reconstruction plates, surgical site will be sutured to end the surgery. During surgery, the amount of blood loss and duration of the operation is measured in both groups. After surgery, mouth opening and post-operative pain is evaluated in multiple time periods. Also the risk of plate exposure and infection during six months of follow-up will be examined.

**Category**

Treatment - Surgery

**2****Description**

intervention2!: In the group using conventional method, CT Scan of axial, coronal and three-dimensional reconstruction views are provided for patients but required treatment plan is determined using only clinical assessment and radiographic evaluation and three-dimensional models will not be manufactured. After routine tests surgery is performed. The amount of mouth opening will be measured before surgery. reconstruction plates are shaped routinely during the surgery and adapted on the bone. Finally, after placement and fixation of reconstruction plates, surgical site will be sutured to end the surgery. During surgery, the amount

of blood loss and duration of the operation is measured. After surgery, mouth opening and post-operative pain is evaluated in multiple time periods. Also the risk of plate exposure and infection during six months of follow-up will be examined.

**Category**

Treatment - Surgery

**Recruitment centers****1****Recruitment center****Name of recruitment center**

Department Of Oral and Maxillofacial Surgery, Faculty Of Dentistry

**Full name of responsible person**

Dr Mohammad Mahdi Nasiri

**Street address**

Department Of Oral and Maxillofacial Surgery, Faculty Of Dentistry

**City**

Yazd

**Sponsors / Funding sources****1****Sponsor****Name of organization / entity**

Vice Chancellor for research of Shahid Sadoghi University of Medical Sciences

**Full name of responsible person**

Dr Amir Hoshang Mehrparvar

**Street address**

Third Floor, Shahid Sadoghi University of Medical Sciences, Bahonar Square

**City**

Yazd

**Grant name****Grant code / Reference number****Is the source of funding the same sponsor organization/entity?**

Yes

**Title of funding source**

Vice Chancellor for research of Shahid Sadoghi University of Medical Sciences

**Proportion provided by this source**

100

**Public or private sector**

empty

**Domestic or foreign origin**

empty

**Category of foreign source of funding**

empty

**Country of origin****Type of organization providing the funding**

empty

**Person responsible for general inquiries****Contact**

**Name of organization / entity**

Shahid Sadoghi University Of Medical Sciences

**Full name of responsible person**

Dr Mohammad Mahdi Nasiri

**Position**

Resident Of Oral & Maxillofacial Surgery

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**Full name of responsible person**

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**Position**

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**Email****Sharing plan****Deidentified Individual Participant Data Set (IPD)**

*empty*

**Study Protocol**

*empty*

**Statistical Analysis Plan**

*empty*

**Informed Consent Form**

*empty*

**Clinical Study Report**

*empty*

**Analytic Code**

*empty*

**Data Dictionary**

*empty*