

Clinical Trial Protocol

Iranian Registry of Clinical Trials

01 Jun 2026

Comparison of the effectiveness of tongue trill, lip trill, straw phonation, and water resistance therapy exercises in patients with primary Muscle Tension Dysphonia

Protocol summary

Study aim

Comparison of the effectiveness of tongue trill, lip trill, straw phonation, and water resistance therapy exercises in patients with primary Muscle Tension Dysphonia

Design

A clinical trial with a control group, parallel groups, double-blind, and randomized on 30 patients

Settings and conduct

The samples will be selected among the clients of the otolaryngology clinic of Amir Alam Hospital and speech therapy clinics of Tehran University of Medical Sciences using the available sampling method

Participants/Inclusion and exclusion criteria

Inclusion criteria: 1) Be female, 2) The age range of 20-50 years, 3) Declare their informed consent to participate in the study, 4) Receiving the diagnosis of primary Muscle tension dysphonia Exclusion criteria: 1) Smoking cigarettes and drinking alcohol; 2) Suffering from hearing loss, hormonal diseases, and neurological diseases; 3) Having a history of laryngeal surgery or laryngeal trauma; 4) Being pregnant; 5) Being menopause; 6) Being menstruating at the time of pre- and post-treatment assessments; 7) Having an upper respiratory tract infection during the assessment days or one week before, and 8) Receiving another voice therapy program at the same time as the study is carried out

Intervention groups

This study has five intervention groups, including four experimental groups and one control group. The experimental groups include lip trill, tongue trill, straw phonation, and water resistance therapy. The control group will receive a vocal hygiene program. All 5 groups will practice their individual treatment plan across 2 weeks with a frequency of two 45-min sessions a week.

Main outcome variables

Maximum Phonation Time; First formant; F1_F0 difference; Cepstral peak prominence; Cepstral peak

prominence-smoothed; Overall Severity of Dysphonia; Frequency and Severity of Vocal Tract Discomfort; Voice Handicap Index

General information

Reason for update

Acronym

IRCT registration information

IRCT registration number: **IRCT20230307057649N1**

Registration date: **2023-09-05, 1402/06/14**

Registration timing: **prospective**

Last update: **2023-09-05, 1402/06/14**

Update count: **0**

Registration date

2023-09-05, 1402/06/14

Registrant information

Name

Roonak Aziz

Name of organization / entity

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Iran (Islamic Republic of)

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Recruitment status

Recruitment complete

Funding source

Expected recruitment start date

2023-09-19, 1402/06/28

Expected recruitment end date

2023-12-19, 1402/09/28

Actual recruitment start date

empty

Actual recruitment end date

empty

Trial completion date

empty

Scientific title

Comparison of the effectiveness of tongue trill, lip trill, straw phonation, and water resistance therapy exercises in patients with primary Muscle Tension Dysphonia

Public title

Comparison of the effectiveness of different semi-occluded vocal tract exercises in patients with primary Muscle Tension Dysphonia

Purpose

Treatment

Inclusion/Exclusion criteria**Inclusion criteria:**

Be female The age range of 20-50 years Announce their informed consent to participate in the study Based on the consensus opinion of an otolaryngologist and an experienced speech therapist in the field of voice disorders and based on the results of voice history taking, larynx observation, palpation examination, auditory-perceptual assessment of voice and self-assessment, receive the diagnosis of primary muscle tension disorder

Exclusion criteria:

Patients who smoke and drink alcohol Patients suffering from hearing loss, hormonal and neurological diseases Patients who have a history of laryngeal surgery and laryngeal trauma Patients who have gone through menopause Patients who receive another voice therapy program at the same time as this study Patients who are pregnant

Age

From **20 years** old to **55 years** old

Gender

Female

Phase

N/A

Groups that have been masked

- Participant
- Outcome assessor

Sample size

Target sample size: **30**

Randomization (investigator's opinion)

Randomized

Randomization description

The studied samples will be randomly divided into five groups to receive one of the five treatments of tongue trill, lip trill, straw phonation, water resistance therapy, and vocal hygiene. Using the rule of random allocation in creating a random sequence and observing the concealment of random allocation is done by Permuted block randomization. For this purpose, 30 people in the sample size of this study are considered as six blocks of five. Each of these six blocks contains five 5 codes of the first to fifth methods. Each block is placed in a separate box containing opaque envelopes containing the codes of five treatment methods, and the blocks are used one

after the other for the lottery. Patients are asked to draw one of the envelopes, take it out and open it and present it to the researcher to be placed in the relevant group

Blinding (investigator's opinion)

Double blinded

Blinding description

In this research, pre-and post-treatment assessments are performed by a speech therapist, and the presentation of therapeutic exercises is by another speech therapist who is the primary researcher of this study. Therefore, the outcome assessor will be blinded to the treatment group allocation, and the principal investigator will be blinded to the assessment results. From the beginning, we will explain to the participants that in this study we intend to compare the effectiveness of several voice therapy methods. We explain to them that these treatment methods have similar goals and are provided to facilitate voice production and solve their voice problems. Next, we explain that the effectiveness of these treatment methods has already been proven and we explain to them about the mechanism of these exercises that this mechanism is implemented using different voice exercises. They are supposed to be randomly placed in one of these treatment groups. Participants with knowledge of the above topics will enter the study if they declare their readiness to participate in this research through the informed consent form. Before the start of the assessment and treatment, the necessary arrangements are made so that the participants do not see each other during the entire period of treatment and assessment. Due to the fact that all participants are assessed and treated individually and will not see other patients, they will not be informed about the allocation of treatment groups.

Placebo

Not used

Assignment

Parallel

Other design features**Secondary Ids**

empty

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

School of Nursing and Midwifery and School of Rehabilitation-Tehran University of Medical Sciences

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Department of Speech Therapy, School of Rehabilitation, Pich Shemiran, Enghelab St., Tehran

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Approval date

2023-04-15, 1402/01/26

Ethics committee reference number

IR.TUMS.FNM.REC.1402.004

Health conditions studied

1

Description of health condition studied

Primary muscle tension dysphonia

ICD-10 code

R49.0

ICD-10 code description

Dysphonia

Primary outcomes

1

Description

Maximum phonation time

Timepoint

Before the start of the intervention and immediately after the completion of four treatment sessions

Method of measurement

Praat Software

2

Description

First Formant

Timepoint

Before the start of the intervention and immediately after the completion of four treatment sessions

Method of measurement

Praat Software

3

Description

The difference between the first formant and the fundamental frequency

Timepoint

Before the start of the intervention and immediately after the completion of four treatment sessions

Method of measurement

Praat Software

4

Description

Cepstral peak prominence

Timepoint

Before the start of the intervention and immediately after the completion of four treatment sessions

Method of measurement

Praat software

5

Description

Cepstral peak prominence-smoothed

Timepoint

Before the start of the intervention and immediately after the completion of four treatment sessions

Method of measurement

Praat software

6

Description

Overall Severity of Dysphonia

Timepoint

Before the start of the intervention and immediately after the completion of four treatment sessions

Method of measurement

Auditory-perceptual evaluation of voice, scale

7

Description

Vocal tract discomfort scale

Timepoint

Before the start of the intervention and immediately after the completion of four treatment sessions

Method of measurement

Self-assessment, The Persian version of the Vocal Tract Discomfort scale

8

Description

Voice Handicap Index

Timepoint

Before the start of the intervention and immediately after the completion of four treatment sessions

Method of measurement

Self-assessment, The Persian version of the VHI questionnaire

Secondary outcomes

empty

Intervention groups

1

Description

First intervention group: The first intervention group will receive voiced tongue trill exercise during four 45-minute sessions over two weeks

Category

Rehabilitation

2

Description

Second intervention group: The second intervention group will receive voiced lip trill exercise during four 45-minute sessions during two weeks

Category

Rehabilitation

3

Description

Third intervention group: The third intervention group will receive straw phonation exercise during four 45-minute sessions during two weeks

Category

Rehabilitation

4

Description

Fourth intervention group: The fourth intervention group will receive water resistance therapy during four 45-minute sessions during two weeks

Category

Rehabilitation

5

Description

Control group: The control group will receive the vocal hygiene program in four 45-minute sessions over two weeks

Category

Lifestyle

Recruitment centers

1

Recruitment center

Name of recruitment center

Amir Alam hospital

Full name of responsible person

Maryam Khodami

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Recruitment center

Name of recruitment center

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Full name of responsible person

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Sponsors / Funding sources

1

Sponsor

Name of organization / entity

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Full name of responsible person

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Grant name

Grant code / Reference number

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

No

Title of funding source

Tehran 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

Tehran University of Medical Sciences

Full name of responsible person

Roonak Aziz

Position

Master Student

Latest degree

Bachelor

Other areas of specialty/work

Speech therapy

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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

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

Analytic Code

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

Data Dictionary

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