

Clinical Trial Protocol

Iranian Registry of Clinical Trials

12 Jun 2026

Survey efficacy of Melissa officinalis on premature ventricular contraction

Protocol summary

Study aim

Determination of the effect of lemon balm on early ventricular heart rate of patients with pulmonary heart disease referred to the clinic of Heshmat Hospital in Rasht

Design

Clinical trials with control group, with parallel groups, blind, randomized

Settings and conduct

The samples were selected from among patients with cardiac palpation who were referred to the Hashamat Heart Center. In order to hide the randomization process from sealed envelopes, which were numbered sequentially, the use and envelope of each person was only after confirmation Eligible eligibility criteria for him and signature of the consent form by the individual will be opened.

Participants/Inclusion and exclusion criteria

Entry requirements: No use of sedative drugs, no drug addiction, no history of herbal medicines allergy, palpitations, ages 18-60, PVC less than 15% Non-compliance: Pvc in the field of cardiac ischemia, patients with valvular disorder, heart failure, thyroid dysfunction, anemia, ACS and cardiac approval, psychosis patients and serious psychiatric illnesses certified by a psychiatrist, endocrinology, pregnancy , Taking betablocker medications, treating hypnosis, and treating anti-anxiety and sedative drugs within 10 days before the study, lack of hypothyroidism.

Intervention groups

The intervention group of 36 patients with a diagnosis of pvc less than 15% based on Holter-rhythm will be included in the study and will be visited monthly by the cardiologist and will be treated in the event of symptoms. These patients will use lemon balm, containing 2 grams (25) of Timen's herbs, for 3 months, twice a day. The control group also includes 36 patients with less than 15% of the pvc and monthly asymptomatic, but are not being treated for medication and will be excluded if symptoms develop.

Main outcome variables

premature ventricular contraction

General information

Reason for update

Acronym

IRCT registration information

IRCT registration number: **IRCT20180205038626N4**

Registration date: **2019-07-30, 1398/05/08**

Registration timing: **registered_while_recruiting**

Last update: **2019-07-30, 1398/05/08**

Update count: **0**

Registration date

2019-07-30, 1398/05/08

Registrant information

Name

Zahra Ahmadnia

Name of organization / entity

Country

Iran (Islamic Republic of)

Phone

+98 13 3361 8177

Email address

zahmadnia@gums.ac.ir

Recruitment status

Recruitment complete

Funding source

Expected recruitment start date

2019-07-23, 1398/05/01

Expected recruitment end date

2020-07-22, 1399/05/01

Actual recruitment start date

empty

Actual recruitment end date

empty

Trial completion date

empty

Scientific title

Survey efficacy of Melissa officinalis on premature ventricular contraction

Public title

Survey efficacy of Melissa officinalis on premature ventricular contraction in Persons with palpitation referring to Dr. Heshmat Rasht Hospital

Purpose

Prevention

Inclusion/Exclusion criteria**Inclusion criteria:**

Tumors use tranquilizers no addiction no history of allergy to herbal medicines palpitations ages 18-60 PVC less than 15%

Exclusion criteria:

Pvc in the field of ischemic heart disease patients with lupus erythematosus heart failure ACS and cardiologist confirmed psychotic patients and serious psychiatric illnesses certified by a psychiatrist endocrine problems pregnancy Treated with hypnosis and treated with anti-anxiety and sedative drugs within 10 days before the study lack of hypothyroidism thyroid dysfunction

Age

From **18 years** old to **60 years** old

Gender

Both

Phase

2

Groups that have been masked

- Data analyser

Sample size

Target sample size: **72**

Randomization (investigator's opinion)

Randomized

Randomization description

Random sequences will be generated using the Random Generator program. Based on the random block method and considering the six blocks, 12 blocks will be generated for 72 patients. After generating the list, each person will be assigned a dedicated code and will be recognized by the code during the study. None of the collaborators participating in the study will be aware of the randomization list, and to use the encryption process to hide the sealed envelopes that are numbered sequentially, the use and envelope of each individual only after confirming the eligibility criteria. Study for him and signature of the consent form by the person will be opened. Registration and random allocation sequences are done by the collaborator of the project methodology and assignment of interventions is done by the collaborator of the plan.

Blinding (investigator's opinion)

Single blinded

Blinding description

In order to hide the randomization process from sealed envelopes, which are numbered sequentially, the use and envelope of each individual will be opened only after confirming the eligible eligibility criteria for him and signing the consent form by the individual.

Placebo

Used

Assignment

Parallel

Other design features**Secondary Ids**

empty

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

Ethics committee of Guilan University of Medical Sciences

Street address

Rasht, Namjoo Ave., Shahid Siadati St., Faced to the 17th Shahrivar Hospital, Old Building, School of Health, University of Technology Research and Technology

City

Rasht

Province

Guilan

Postal code

41446-66949

Approval date

2019-07-21, 1398/04/30

Ethics committee reference number

IR.GUMS.REC.1398.178

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

premature ventricular contraction

ICD-10 code

I49.3

ICD-10 code description

Ventricular premature depolarization

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

premature ventricular contraction

Timepoint

At the end of the 3-month period, both rituals and echo cancers will be performed and the amount of pvc reduction will be evaluated by the cardiologist.

Method of measurement

Echocardiography, Holler rhythm and ECG

Secondary outcomes

empty

Intervention groups

1

Description

Intervention group: The intervention group of 36 patients with a diagnosis of pvc less than 15% based on Holter rhythm will be included in the study and will be visited monthly by a cardiologist and will be excluded from the study in the event of symptoms of shortness of breath, cough, chest pain, pulmonary heart disease. They are under medical treatment. These patients will use lemon balm for 3 months twice a day.

Category

Other

2

Description

Control group: The control group also includes 36 patients with less than 15% of the pvc and monthly asymptomatic, but are not being treated for medication and will be excluded if symptoms develop.

Category

Other

Recruitment centers

1

Recruitment center

Name of recruitment center

Dr. Heshmat Rasht Hospital

Full name of responsible person

Arsalan Salari

Street address

Dr. Heshmat Rasht Hospital -Masala Square, Baniyan St.

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4193955588

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salari@gums.ac.ir

Sponsors / Funding sources

1

Sponsor

Name of organization / entity

Rasht University of Medical Sciences

Full name of responsible person

Shadman Nemati

Street address

Namjoo Ave., Shahid Siadati St., Rasht., Technology & Research Vice-chancellor of University

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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

Rasht 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

Rasht University of Medical Sciences

Full name of responsible person

Arsalan Salari

Position

Associate professor

Latest degree

Specialist

Other areas of specialty/work

Cardiology

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Person responsible for scientific inquiries

Contact

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Person responsible for updating data

Contact

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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