

Clinical Trial Protocol

Iranian Registry of Clinical Trials

10 Jun 2026

Comparison of the effect of topical solution of Finasteride 0/2% versus topical solution of Minoxidil 5% in the treatment of Androgenic Alopecia in men, A Randomized, Double-Blind, Clinical Trial

Protocol summary

Study aim

Determining and comparing the effect of two drugs, topical solution of Finasteride 0/2% and topical solution of Minoxidil 5% in the treatment of Androgenetic Alopecia in men.

Design

Clinical trial with control group, with parallel groups, Double blind, Random, phase 2 on 200 patients, for randomization, method alternate was used.

Settings and conduct

Patients are randomly divided into two groups of 100 people. Receive as one among bottles A and B. After daily use according to the instructions, they are monitored on a monthly basis and a checklist for each patient is completed at each visit. This process lasts up to 6 months. Monthly visit is done in the Dermatology Clinic of Imam Reza Hospital in Ardabil on sundays and wednesdays every week. During this study, neither the patient nor the attending physician was aware of the contents of the bottle prescribed to the patient and at the end of the study, after collecting and analyzing the data, the nature of bottles A and B will be determined.

Participants/Inclusion and exclusion criteria

If there is more than a week between medications, the patient will be excluded from the study. Patients under study during these six months should not use any other drugs (traditional and industrial), either topically or systemically because If used, they will be excluded from the study.

Intervention groups

Patients are randomly divided into two groups of 100 people. To a group (intervention) bottle containing 25 cc of 0/2% topical Finasteride solution and the second group (control) is given 25 cc of 5% Minoxidil topical solution. Patients should be examined monthly.

Main outcome variables

The number of hairs plucked in the Pull test; Active areas

for Alopecia

General information

Reason for update

Change in the percentage of Finasteride (from 2% to 0.2%) due to a mistake in entering the initial information.

Acronym

IRCT registration information

IRCT registration number: **IRCT20191023045213N1**

Registration date: **2020-08-19, 1399/05/29**

Registration timing: **prospective**

Last update: **2020-09-01, 1399/06/11**

Update count: **1**

Registration date

2020-08-19, 1399/05/29

Registrant information

Name

Amir Sadrinia

Name of organization / entity

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Iran (Islamic Republic of)

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+98 21 4472 1657

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Recruitment status

Recruitment complete

Funding source

Expected recruitment start date

2020-09-22, 1399/07/01

Expected recruitment end date

2020-10-20, 1399/07/29

Actual recruitment start date

empty

Actual recruitment end date

empty

Trial completion date

empty

Scientific title

Comparison of the effect of topical solution of Finasteride 0/2% versus topical solution of Minoxidil 5% in the treatment of Androgenic Alopecia in men, A Randomized, Double-Blind, Clinical Trial

Public title

The effect of Finasteride solution on male hair loss

Purpose

Treatment

Inclusion/Exclusion criteria**Inclusion criteria:**

Volunteer to participate in the study

Exclusion criteria:**Age**

No age limit

Gender

Male

Phase

3

Groups that have been masked

- Participant
- Care provider
- Investigator
- Outcome assessor

Sample size

Target sample size: **200**

Randomization (investigator's opinion)

Randomized

Randomization description

In this study, there will be two types of dark bottles, one named A and the other B. In one type of bottle 25 ml of Finasteride 0/2% topical solution and in the second type 25 ml of Minoxidil 5% topical solution are poured. The patients were randomly divided into two groups of 100 each and In the first stage, they are subjected to a Pull test, which is an important test to determine the active sites of Alopecia and In it 60-70 pieces of patient hair are grabbed and pulled hairs are counted and If the number of plucked hairs is more than 6-7, the test is positive and the area of the head is considered as the active site for Alopecia and its subsequent monthly examination. In subsequent monthly examinations, those points on the head will be tested for loss. Then, one group is given bottle A and the other group is given bottle B. The patient should apply 15 drops of these solutions on his head every 12 hours and at the end of each month, go to the clinic for a checkup. In the monthly examination, patients also undergo a pull test and the number of hairs plucked in the test is recorded, until the end of the sixth month, which is the end of treatment. If the patient runs out of medicine during this time, he will receive a bottle with the same name as the first bottle, and if he forgets to use his medication on time, he should use his medication immediately after being reminded, but if there is a gap of more than a week between taking the

drug the patient will be excluded from the study. Patients under study should not use any other drugs (traditional and industrial) during these six months, either topically or systemically as they will be excluded if the study is used, If used, they will be excluded from the study.

Blinding (investigator's opinion)

Double blinded

Blinding description

During this study, a dermatologist as the head of the research group in the clinic, after explaining the working method for patients and possible problems during the research period for patients, if he volunteers, he puts them in the study. In order to start treatment, they enter group A or B as one of the volunteer patients. After assigning the checklist to each patient (according to the entered group), the patient receives bottles containing the treatment solution (A or B) from the head of the clinic according to his checklist. During this process, neither the specialist physician nor the clinic manager nor the patient don't knows the contents of the therapeutic bottle and which group of bottles are Finasteride or Minoxidil. Therapeutic bottles are made by a pharmacologist in the university laboratory and poured into bottles and then transported to the clinic of the study. The pharmacologist is the only person who knows the contents of the bottles and during the study and analysis of the data has nothing to do with the dermatologist and the head of clinic and the analyzer and patients. After the end of sampling and the end of the 6-month treatment period, by analyzing the data, the analyzer presents the information in the form of an initial grouping, and at the end of the analysis, the nature of the bottles will be determined by a pharmacologist.

Placebo

Not used

Assignment

Parallel

Other design features**Secondary Ids**

empty

Ethics committees

1

Ethics committee**Name of ethics committee**

Ethics committee of Ardebil University of Medical Sciences

Street address

Daneshgah Ave

City

Ardebil

Province

Ardabil

Postal code

5618985991

Approval date

2020-06-01, 1399/03/12

Ethics committee reference number

Health conditions studied

1

Description of health condition studied

Androgenic Alopecia

ICD-10 code

L64

ICD-10 code description

Androgenic alopecia

Primary outcomes

1

Description

Pull test score

Timepoint

Monthly

Method of measurement

Physical exam

Secondary outcomes

empty

Intervention groups

1

Description

Control group: After receiving a topical solution of Minoxidil and taking the drug every 12 hours 15 mg, they are included in the study and examined monthly.

Category

Treatment - Drugs

2

Description

Intervention group: After receiving a topical solution of Finasteride and taking it according to Minoxidil, they are included in the study and examined monthly

Category

Treatment - Drugs

Recruitment centers

1

Recruitment center

Name of recruitment center

Ardebil's Imam Reza hospital

Full name of responsible person

Majid Rostami Moghadam

Street address

Basij Sq

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Web page address

Sponsors / Funding sources

1

Sponsor

Name of organization / entity

Ardebil University of Medical Sciences

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?

Yes

Title of funding source

Ardebil 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

Ardebil University of Medical Sciences

Full name of responsible person

Majid Rostami Moghadam

Position

Professor

Latest degree

Specialist

Other areas of specialty/work

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Person responsible for updating data**Contact****Name of organization / entity**

Ardabil University of Medical Sciences

Full name of responsible person

Amir Sadrinia

Position

Consultant

Latest degree

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General Practitioner

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

Yes - There is a plan to make this available

Study Protocol

No - There is not a plan to make this available

Statistical Analysis Plan

Not applicable

Informed Consent Form

No - There is not a plan to make this available

Clinical Study Report

Not applicable

Analytic Code

Not applicable

Data Dictionary

Not applicable

Title and more details about the data/document

All data is potentially shareable after unidentified individuals.

When the data will become available and for how long

Access started from 1401

To whom data/document is available

Researchers working in academic and scientific institutions.

Under which criteria data/document could be used

If the purpose of access is to investigate possible bugs in the study.

From where data/document is obtainable

Amir Sadrinia Email address: sadriamir73@gmail.com

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

As soon as the email is sent to the mentioned address, the information will be sent with the confirmation of the purpose.

Comments