

Clinical Trial Protocol

Iranian Registry of Clinical Trials

17 Jun 2026

Comparative study on the effectiveness and safety of tamsulosin Oral-Controlled Absorption System (OCAS®) with tamsulosin Modified Release (MR) and placebo in improving lower urinary tract symptoms (LUTS) in men with benign prostatic hyperplasia (BPH)

Protocol summary

Study aim

Compararison of the efficacy and safety of tamsulosin Oral-Controlled Absorption System (OCAS®) with tamsulosin Modified Release (MR) and placebo in improving lower urinary tract symptoms (LUTS) in men with benign prostatic hyperplasia (BPH)

Design

Phase 3 clinical trial with control group, with parallel groups, double-blind, randomized with random allocation method, with a sample size of 260 patients

Settings and conduct

Medications are separately packed in 4 groups from 1 to 4 by one of the co-workers and delivered to the main researcher. So that neither patients nor the main researcher who prescribes the medication do not know the type of drug they receive or is prescribed. The location of the study is the Urology Clinic of Razi Educational and Research Hospital. Drugs are randomly distributed among eligible patients after necessary examinations.

Participants/Inclusion and exclusion criteria

Patients complained of lower urinary tract symptoms due to benign prostatic hyperplasia with age of at least 40 years, Prostate-Specific Antigen (PSA) in age-specific range and score of 8 or greater in the international prostate symptom score (IPSS) questionnaire are included and patient with renal, cardiovascular and liver disorders are not included.

Intervention groups

Group 1: Tamsulosin Capsule 0.4 mg (MR) with brand name Portral made by Tasnim Pharmaceutical Co., daily oral administration for 12 weeks; Group 2: Talsulosin 0.4 mg (OCAS) with brand name Protral OPAS, made by Tasnim Pharmaceutical Co., daily oral administration for 12 weeks; Group 3: Tamsulosin 0.4 mg (OCAS) with brand name Omnic OCAS, daily oral administration for 12

weeks; Group 4: Placebo, made by Tasnim Pharmaceutical Co., daily oral administration for 12 weeks;

Main outcome variables

International prostate symptom score (IPSS) questionnaire

General information

Reason for update

Acronym

IRCT registration information

IRCT registration number: **IRCT20100817004582N8**

Registration date: **2018-02-17, 1396/11/28**

Registration timing: **registered_while_recruiting**

Last update: **2018-02-17, 1396/11/28**

Update count: **0**

Registration date

2018-02-17, 1396/11/28

Registrant information

Name

Ali Hamidi Madani

Name of organization / entity

Urology Research Center, Guilan University of Medical Sciences

Country

Iran (Islamic Republic of)

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Recruitment status

Recruitment complete

Funding source

Expected recruitment start date

2018-01-21, 1396/11/01

Expected recruitment end date

2019-01-21, 1397/11/01

Actual recruitment start date

empty

Actual recruitment end date

empty

Trial completion date

empty

Scientific title

Comparative study on the effectiveness and safety of tamsulosin Oral-Controlled Absorption System (OCAS®) with tamsulosin Modified Release (MR) and placebo in improving lower urinary tract symptoms (LUTS) in men with benign prostatic hyperplasia (BPH)

Public title

Comparison of the efficacy and safety of tamsulosin OCAS with tamsulosin MR and placebo in improving lower urinary tract symptoms

Purpose

Treatment

Inclusion/Exclusion criteria**Inclusion criteria:**

Patients who are at least 40 years of age. Prostate-Specific Antigen (PSA) score should be at age-specific range. International Prostate Symptom Score (IPSS) should be greater than or equal to 8. The maximum urine flow (Qmax) should be in the range of greater than and/or equal to 4 ml/s and smaller and/or equal to 15 ml/s. The urine residual should be below 120 ml.

Exclusion criteria:

History of prostatectomy Severe liver dysfunction Severe renal dysfunction Severe cardiovascular dysfunction History of syncope. Other conditions that can be causing voiding dysfunction, such as neurogenic bladder, bladder or urinary stones, frequent urinary tract infection, bladder cancer, prostate cancer, urethral stenosis, and large diverticulum bladder. History of allergy to tamsulosin History of recent retention and nocturnal polyuria Patients with benign prostatic hyperplasia (BPH) who are candidates for surgery on the basis of clinical and paraclinical symptoms. History of use of 5 α -reductase inhibitors in the last 3 months History of taking diuretics and hypnotic tablets Use of other therapies such as other alpha blockers or herbal extracts in the last month Use of other drugs, such as alpha-agonists, cholinergic or anticholinergic drugs that may affect the effects of tamsulosin pharmacodynamics.

Age

From 40 years old

Gender

Male

Phase

4

Groups that have been masked

- Participant
- Investigator

Sample size

Target sample size: 260

Randomization (investigator's opinion)

Randomized

Randomization description

Patients are randomly assigned through random allocation method with block size 8 to 4 groups.

Blinding (investigator's opinion)

Double blinded

Blinding description

Medications are separately packaged in 4 groups of 1 to 4 by one of the co-workers and delivered to the main researcher.

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

Opposite of 17 Shahrivar Hospital, Shahid Siadati Street, Namjoo Street

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Rasht

Province

Guilan

Postal code

41446-66949

Approval date

2017-11-11, 1396/08/20

Ethics committee reference number

IR.GUMS.REC.1396.318

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

Lower Urinary Tract Symptoms (LUTS)

ICD-10 code

N40.1

ICD-10 code description

Enlarged prostate with lower urinary tract symptoms

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

The International Prostate Symptom Score (IPSS) Questionnaire

Timepoint

The beginning of the study (before the intervention) and 6 and 12 weeks after the start of the intervention

Method of measurement

The International Prostate Symptom Score (IPSS) Questionnaire

Secondary outcomes

1

Description

Side effects

Timepoint

The beginning of the study (before the intervention) and 6 and 12 weeks after the start of the intervention

Method of measurement

Ask the patient

Intervention groups

1

Description

Intervention group 1: Tamsulosin capsule 0.4 mg (MR) with brand name Portral, manufacturing by Tasnim Pharmaceutical Co., daily for 12 weeks orally

Category

Treatment - Drugs

2

Description

Intervention group 2: Tamsulosin Tablets 0.4 mg (OCAS) with brand name Protral OPAS, manufacturing by Tasnim Pharmaceutical Co., daily for 12 weeks orally

Category

Treatment - Drugs

3

Description

Intervention group 3: Tamsulosin Tablets 0.4 mg (OCAS) with brand name Omnic OCAS, daily for 12 weeks orally

Category

Treatment - Drugs

4

Description

Control group: Placebo, manufacturing by Tasnim Pharmaceutical Co., daily for 12 weeks orally

Category

Placebo

Recruitment centers

1

Recruitment center

Name of recruitment center

Razi Hospital

Full name of responsible person

Ali Hamidi Madani

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Razi Hospital, Sardar Jangal Street

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Sponsors / Funding sources

1

Sponsor

Name of organization / entity

Urology Research Centre, Guilan University of Medical Sciences

Full name of responsible person

Head of Urology Research Centre, Dr. Siavsh Falahatkar

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Urology Research Centre, Razi Hospital, Sardar Jangal St.

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Grant name

10506

Grant code / Reference number

32

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

Yes

Title of funding source

Urology Research Centre, Guilan University of Medical Sciences

Proportion provided by this source

20

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

2**Sponsor****Name of organization / entity**

Tasnim Pharmaceutical Co.

Full name of responsible person

Dr. Abbas Movahednia

Street address

Unit 6, 2nd floor, No. 11, 1st Alley, Beginning at Gandhi Street

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Email

tasnimpharam@yahoo.com

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

Yes

Title of funding source

Tasnim Pharmaceutical Co.

Proportion provided by this source

80

Public or private sector

Private

Domestic or foreign origin

Domestic

Category of foreign source of funding*empty***Country of origin****Type of organization providing the funding**

Industry

Person responsible for general inquiries**Contact****Name of organization / entity**

Urology Research Center, Guilan University of Medical Sciences

Full name of responsible person

Kourosh Mojtavavi

Position

Medical student

Latest degree

Medical doctor

Other areas of specialty/work

Urology

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Person responsible for scientific inquiries**Contact****Name of organization / entity**

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Full name of responsible person

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Position

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Specialist

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

Urology Research Center, Guilan University of Medical Sciences

Full name of responsible person

Samaneh Esmaeili

Position

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Latest degree

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Other areas of specialty/work

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samaneh_815@yahoo.com

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

No - There is not a plan to make this available

Justification/reason for indecision/not sharing IPD

It is not ethically possible to release patient data. The questionnaires are kept at the Urology Research Center of Guilan University of Medical Sciences and only if necessary, natural and legal persons with legal permission will be allowed to examine them.

Study Protocol

Yes - There is a plan to make this available

Statistical Analysis Plan

Yes - There is a plan to make this available

Informed Consent Form

No - There is not a plan to make this available

Clinical Study Report

Yes - There is a plan to make this available

Analytic Code

Yes - There is a plan to make this available

Data Dictionary

Not applicable

Title and more details about the data/document

All patient data will be reported after a statistical analysis in the form of the results of a scientific study.

When the data will become available and for how long

Start the access period after printing the results in a scientific article

To whom data/document is available

All researchers in Iran who are affiliated with one of the academic or research institutes approved by the Ministry of Health and Medical Education.

Under which criteria data/document could be used

In case of doing Cohort studies or multicentre studies by other eligible researchers

From where data/document is obtainable

Dr. Ali Hamidi Madani, executor of project and Dr. Siavash Falahtakar, Head of Urology Research Center
Address: Urology Research Center, Razi Hospital, Rasht
Phone: 013- 33525259

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

Submitting a written request for the head of the Urology Research Center, stating the request to the project executive other colleagues and getting their approval, writing a written opinion to the applicant, taking the moral obligation of the applicant, sending the data and documentation to the applicant

Comments