

Clinical Trial Protocol

Iranian Registry of Clinical Trials

13 Jun 2026

Evaluation of efficacy of Maa-al-shaeer (An Iranian medicine product based on barley seeds) in preventing and reducing the severity of radiotherapy-induced dysuria in prostate cancer patients; a double-blind randomized clinical trial

Protocol summary

Study aim

Evaluation of the efficacy of Maa-al-shaeer in preventing and reducing the severity of radiotherapy-induced dysuria in prostate cancer

Design

A controlled, parallel-group, double-blind, randomized, phase 3 clinical trial on 70 patients. Randomization is done using R software version 4.1.2 and the randomization list generated using R software will be used.

Settings and conduct

Prostate cancer patients candidates for radiotherapy (in the radiotherapy center of Shohada-e-Tajrish Hospital), without the exclusion criteria, will be included in the study and assigned to the intervention or control group randomly. The study is double-blind, and patients, healthcare providers, researchers, and analysts are all blinded.

Participants/Inclusion and exclusion criteria

Inclusion criteria: 1. Over 18 years old 2. Prostate cancer 3. Candidate for radiation therapy 4. Consciousness
Criteria for not entering the study: 1. Unconsciousness or unable to report the dysuria 2. Urine catheter 3. Urine analysis test with infection evidence 4. Diabetes Mellitus 5. Dysuria 6. History of drug sensitivity 7. Metastasis 8. Chemotherapy at the same time 9. Hematuria 10. Severe cardiovascular, pulmonary, hepatic, and kidney disease needs to be treated 11. Not participating in the study

Intervention groups

The participants, twice a day, each time, a measure (equivalent to 7.5 cc) of powder (Maa-al-shaeer in the intervention group or placebo in the control group) dissolve in a glass of water and consume it. The duration of using the drug or placebo is 4 weeks from the beginning of radiotherapy.

Main outcome variables

Dysuria, Quality of life, The number of phenazopyridine tablets which was taken, Treatment satisfaction (secondary outcome)

General information

Reason for update

Acronym

IRCT registration information

IRCT registration number: **IRCT20190608043840N1**

Registration date: **2023-01-22, 1401/11/02**

Registration timing: **prospective**

Last update: **2023-01-22, 1401/11/02**

Update count: **0**

Registration date

2023-01-22, 1401/11/02

Registrant information

Name

Vida Nazari

Name of organization / entity

Country

Iran (Islamic Republic of)

Phone

+98 21 2230 3841

Email address

vidanazary@sbmu.ac.ir

Recruitment status

Recruitment complete

Funding source

Expected recruitment start date

2023-02-01, 1401/11/12

Expected recruitment end date

2023-07-01, 1402/04/10

Actual recruitment start date

empty

Actual recruitment end date

empty

Trial completion date

empty

Scientific title

Evaluation of efficacy of Maa-al-shaeer (An Iranian medicine product based on barley seeds) in preventing and reducing the severity of radiotherapy-induced dysuria in prostate cancer patients; a double-blind randomized clinical trial

Public title

Evaluation of the efficacy of Maa-al-shaeer (product based on barley seeds) on radiotherapy induced dysuria

Purpose

Prevention

Inclusion/Exclusion criteria**Inclusion criteria:**

Over 18 years old Prostate cancer Candidate for radiation therapy (at the Radiotherapy Center of Shohada-e-Tajrish Hospital) Consciousness

Exclusion criteria:

Unconsciousness or unable to report dysuria Urinary catheter Urinary tract infection Diabetes Mellitus Dysuria A history of drug sensitivity Metastasis Chemotherapy at the same time Hematuria Severe Cardiovascular, Pulmonary, Hepatic and Kidney diseases that need some medical treatments Not participating in the study

Age

From **18 years** old

Gender

Male

Phase

3

Groups that have been masked

- Participant
- Care provider
- Investigator
- Outcome assessor
- Data analyser

Sample size

Target sample size: **70**

Randomization (investigator's opinion)

Randomized

Randomization description

Simple Randomization. Randomization is done using R software version 4.1.2, and we will use the randomization list generated using R software. In order to conduct this study, first, using the available method, patients who meet the conditions for entering the study, after completing the consent form, will enter the study until the number of calculated samples is complete. In the next step, in order to homogenize and control the background and confounding variables in the intervention and control groups, the block randomization method will be used to assign patients to two groups. For this purpose and according to the calculated sample size, 5 blocks of 14 blocks will be generated using R software.

In each block, the interventions are randomly assigned to people with a ratio of 1:1. Since the shape of the drug and the placebo are completely similar and the labels on the cans of the interventions (drug or placebo) are prepared by a statistician (using R software) and stucked in the pharmaceutical laboratory, so, the participants and the researcher are all blind to it.

Blinding (investigator's opinion)

Double blinded

Blinding description

Double Blind. In this study, the barley product is prepared in the form of powder and is packed in a can and coded after putting a label. Placebo is also prepared with the same particle size and color as the drug in the same packaging as the drug and is coded. Therefore, until the encoding, it is not possible for the participants, principle investigator, healthcare providers, data collectors, outcome assessors, manuscript writers to distinguish the drug from the placebo.

Placebo

Used

Assignment

Parallel

Other design features**Secondary Ids**

empty

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

Ethics Committee of Shahid Beheshti University of Medical Sciences

Street address

Shahid Beheshti University of Medical Sciences, Sh. Aarabi Ave., Yaman Ave., Velenjak

City

Tehran

Province

Tehran

Postal code

1985717443

Approval date

2022-09-12, 1401/06/21

Ethics committee reference number

IR.SBMU.RETECH.REC.1401.367

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

Prostate Cancer, Radiotherapy-induced dysuria, Urogenital radiotherapy complications

ICD-10 code

Malignant

ICD-10 code description

C61

Primary outcomes

1

Description

Dysuria

Timepoint

Before intervention and 2, 4, 6, 8 weeks after intervention

Method of measurement

IPSS (International prostate symptom score) questionnaire

Secondary outcomes

1

Description

Quality of Life

Timepoint

Quality of life score before the intervention (week 0) 4 and 8 weeks after the intervention

Method of measurement

EORTC QLQ-PR25 Questionnaire (European Organization for Research and Treatment of Cancer Quality of Life Questionnaire for Prostate Cancer)

2

Description

The number of phenazopyridine tablets which was used

Timepoint

2, 4, 6 and 8 weeks after the intervention

Method of measurement

Asking the participants

3

Description

Treatment satisfaction

Timepoint

8 weeks after the intervention

Method of measurement

It is measured by asking 1 Likert scale question (very high, high, medium, low and very low).

Intervention groups

1

Description

Intervention group: Twice a day, every 12 hours, one measure of "Maa-al-shaeer" (malted barley powder which was prepared by the school of Traditional Medicine of Shahid Beheshti University of Medical Sciences) is dissolved in a glass of lukewarm water and consumed after meals. Each measure of powder is equivalent to 7.5 cc of powder and the intervention period is four weeks from the start of radiation therapy and the follow-up period is four weeks (without intervention).

Category

Prevention

2

Description

Control group: Twice a day, every 12 hours, one measure of placebo powder (which was prepared by the school of Traditional Medicine of Shahid Beheshti University of Medical Sciences) is dissolved in a glass of lukewarm water and consumed after meals. Each measure of powder is equivalent to 7.5 cc of powder and the intervention period is four weeks from the start of radiation therapy and the follow-up period is four weeks (without intervention).

Category

Prevention

Recruitment centers

1

Recruitment center

Name of recruitment center

Shohada Tajrish Hospital⁹⁹

Full name of responsible person

Vida Nazari

Street address

Shohada of Tajrish Hospital, Shahrdari st.,Tajrish sq.

City

Tehran

Province

Tehran

Postal code

1989934148

Phone

+98 21 25719

Email

Pr_shohada@sbmu.ac.ir

Sponsors / Funding sources

1

Sponsor

Name of organization / entity

Shahid Beheshti University of Medical Sciences

Full name of responsible person

Dr. Afshin Zarghi

Street address

7th Floor, Bldg No.2 SBUMS, Arabi AveTehran Prov7th Floor, Bldg No.2 SBUMS, Arabi Ave, Velenjak, Tehran

City

Tehran

Province

Tehran

Postal code

19839 69411

Phone

+98 21 2243 9781

Email

Mpajouhesh@sbmu.ac.ir

Grant name

Grant code / Reference number

Is the source of funding the same sponsor

organization/entity?

Yes

Title of funding source

Shahid Beheshti 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

University of Medical Sciences, No. 8, Shams Alley, In front of Tavanir Ave., Vali-Asr Ave., Tehran

City

Tehran

Province

Tehran

Postal code

1516745811

Phone

+98 21 8877 3521

Fax

+98 21 8879 5008

Email

vidanazary@sbmu.ac.ir

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

Shahid Beheshti University of Medical Sciences

Full name of responsible person

Vida Nazari

Position

Phd Student

Latest degree

Medical doctor

Other areas of specialty/work

Traditional Medicine

Street address

Faculty of Traditional Medicine, Shahid Beheshti University of Medical Sciences, No. 8, Shams Alley, In front of Tavanir Ave., Vali-Asr Ave., Tehran

City

Tehran

Province

Tehran

Postal code

1516745811

Phone

+98 21 8877 3521

Fax

+98 21 8879 5008

Email

vidanazary@sbmu.ac.ir

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

Shahid Beheshti University of Medical Sciences

Full name of responsible person

Vida Nazari

Position

Phd Student

Latest degree

Medical doctor

Other areas of specialty/work

Traditional Medicine

Street address

Faculty of Traditional Medicine, Shahid Beheshti University of Medical Sciences, No. 8, Shams Alley, In front of Tavanir Ave., Vali-Asr Ave., Tehran

City

تهران

Province

Tehran

Postal code

1516745811

Phone

+98 21 8877 3521

Fax

+98 21 8879 5008

Email

Vidanazary@sbmu.ac.ir

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

Shahid Beheshti University of Medical Sciences

Full name of responsible person

Vida Nazari

Position

Phd Student

Latest degree

Medical doctor

Other areas of specialty/work

Traditional Medicine

Street address

Faculty of Traditional Medicine, Shahid Beheshti

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

The study has not yet started, after the study is completed a decision will be made about sharing the participants' non-identifiable personal data.

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