

Clinical Trial Protocol

Iranian Registry of Clinical Trials

18 Jun 2026

The effect of the motion exercise program on the quality of life in prostate cancer survivors

Protocol summary

Study aim

Determine the effect of exercise program on the quality of life of survivors of prostate cancer

Design

In this study, survivors of prostate cancer who had a medical record in the center of radiotherapy at Imam Khomeini Hospital in Tehran and had criteria for entering the study, were the sample of the study. Patients with selected entry criteria were selected by random sampling method. A random sample of 80 people entered the study and randomly divided into intervention and control groups. To conceal the intervention group, each participant had a code is assigned.

Settings and conduct

Patients in the intervention group were divided into 5 groups of 8, according to their area of living and their convenient access to sports facilities in the park near the place of residence. At first, the benefits of physical activity and how the exercise program was performed during the research period by the researcher was taught to the participants in each of the 5 groups and a exercise program was provided to them. Regarding the coordination with the municipality of Tehran, the patients in the intervention group during the study, which lasted for 12 weeks, performed one weekly session in a group under the supervision of a researcher in the park's physical environment and three sessions per week individually. Aerobic walking was performed in walking style with mild to moderate intensity (based on patient tolerance). Resistive exercise movements were performed for muscle strength with mild to moderate intensity (based on patient tolerance) for each large muscle group twice a week, once grouped. Stretching exercises for pulling of the main muscles and tendons were also done twice a week in a single session, individually and one session. In addition, patients with urinary incontinence were trained and recommended for pelvic floor exercise. Which performs these movements daily. Questionnaires related to quality of life were

completed before and after the intervention in both control and intervention groups.

Participants/Inclusion and exclusion criteria

Inclusion criteria: Patients diagnosed with prostate cancer and completed active treatment within 3 to 12 months The absence of metastasis to other locations The absence of neurological, cardiovascular, respiratory or any disorder that prevents physical activity. At least 45 years of age exclusion criteria: Changes in health condition that prevent physical activity

Intervention groups

In this study there is an intervention group that carries out exercise exercises. And there is a control group that there will be no action or intervention in this group.

Main outcome variables

quality of life

General information

Reason for update

Acronym

IRCT registration information

IRCT registration number: **IRCT20180113038347N1**

Registration date: **2018-01-23, 1396/11/03**

Registration timing: **retrospective**

Last update: **2018-01-23, 1396/11/03**

Update count: **0**

Registration date

2018-01-23, 1396/11/03

Registrant information

Name

Abbas Mardani

Name of organization / entity

Country

Iran (Islamic Republic of)

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Recruitment status

Recruitment complete

Funding source**Expected recruitment start date**

2017-05-21, 1396/02/31

Expected recruitment end date

2017-10-21, 1396/07/29

Actual recruitment start date

2017-05-21, 1396/02/31

Actual recruitment end date

2017-12-20, 1396/09/29

Trial completion date

empty

Scientific title

The effect of the motion exercise program on the quality of life in prostate cancer survivors

Public title

"The effect of the motion exercise program on the quality of life in prostate cancer survivors"

Purpose

Education/Guidance

Inclusion/Exclusion criteria**Inclusion criteria:**

Patients diagnosed with prostate cancer and completed active treatment within the past 3 to 12 months Lack of metastasis The absence of neurological, cardiovascular, respiratory or any other problems that prevent physical activity At least 45 years of age Conscious consent to participate in the study Residence in Tehran

Exclusion criteria:

Absent more than three sessions in group exercise sessions Changes in health condition that prevent physical activity

Age

From 45 years old

Gender

Male

Phase

N/A

Groups that have been masked

No information

Sample size

Target sample size: 80

Actual sample size reached: 71

Randomization (investigator's opinion)

Randomized

Randomization description

The method of sampling in this research is simple random sampling using random numbers table. In this way, patients with prostate cancer who have criteria for entering the study are selected, then each patient is given a code and using the random numbers table, the first code is equal or smaller than the number of patients eligible for the study Enter the control group and one in each patient according to the code assigned to the intervention and control group, and to some extent in the random numbers table we go to each of the intervention

and control groups of 40 patients.

Blinding (investigator's opinion)

Not blinded

Blinding description**Placebo**

Not used

Assignment

Other

Other design features**Secondary Ids**

empty

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

کمیته اخلاق دانشگاه علوم پزشکی تهران

Street address

Tehran University of Medical Sciences, Nursing and Midwifery Faculty, Tehran, Iran

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1419733171

Approval date

2017-05-02, 1396/02/12

Ethics committee reference number

IR.TUMS.FNM.REC.1396.2209

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

Prostate Cancer Survivors

ICD-10 code

C61

ICD-10 code description

Malignant neoplasm of prostate

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

Quality of Life

Timepoint

At the beginning of the study (before the intervention) and after the end of the intervention (duration of intervention 12 weeks)

Method of measurement

Using EORTC QLQ-C30 Quality of Life Questionnaire. V 3 and EORTC QLQ-PR25 belonging to the European Cancer Research & Treatment Organization. These two complementary questionnaires are also used to measure the quality of life of patients with prostate cancer.

Secondary outcomes

empty

Intervention groups

1

Description

Intervention group: Patients in the intervention group were divided into 5 groups of 8, according to their area of living and their convenient access to sports facilities in the park near the place of residence. At first, the benefits of physical activity and how the exercise program was performed during the research period by the researcher was taught to the participants in each of the 5 groups and a motor exercise program was provided to them. Regarding the coordination with the municipality of Tehran, the patients in the intervention group during the study, which lasted for 12 weeks, performed one weekly session in a group under the supervision of a researcher in the park's physical environment and three sessions per week individually. Aerobic walking was performed in walking style with moderate to moderate intensity (based on patient tolerance). Resistive exercise movements were performed for muscle strength with moderate to moderate intensity (based on patient tolerance) for each large muscle group twice a week, once grouped. Stretching exercises for pulling the main muscles and tendons were also performed twice a week in a single session, individually and one session. In addition, patients with urinary incontinence were trained and recommended for pelvic floor exercise. Which performs these movements daily.

Category

N/A

2

Description

Control group: No specific action was taken in patients in the control group.

Category

N/A

Recruitment centers

1

Recruitment center

Name of recruitment center

بیمارستان آموزشی امام خمینی شهرتهران وابسته به دانشگاه علوم پزشکی تهران

Full name of responsible person

عباس مردانی

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Sponsors / Funding sources

1

Sponsor

Name of organization / entity

Deputy of Research of Tehran University of Medical Sciences

Full name of responsible person

Shadan Pedram Razi

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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

Deputy of Research of 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

Abbas mardani

Position

Student

Latest degree

Bachelor

Other areas of specialty/work

Nursery

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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

To protect the privacy of the participants in the study

Study Protocol

No - There is not a plan to make this available

Statistical Analysis Plan

No - There is not a plan to make this available

Informed Consent Form

No - There is not a plan to make this available

Clinical Study Report

No - There is not a plan to make this available

Analytic Code

No - There is not a plan to make this available

Data Dictionary

No - There is not a plan to make this available