

Clinical Trial Protocol

Iranian Registry of Clinical Trials

29 Aug 2026

The impact of a results-based motivating system on population levels of the non-communicable diseases risk factors in Iran: A field trial study

Protocol summary

Study aim

Study the impact of a results-based motivating system on population levels of the non-communicable diseases risk factors in Iran: A field trial study

Design

First phase: at this stage, the basic information is collected based on the STEP questionnaire (with biochemical and physical measurements), at which point all four groups will be included in the study. The second phase: by using regular review studies, the best evidence and best practices will be obtained for effective interventions, and then we will train it with health experts and carers to use the methods. Third phase: in this phase, an operational plan with the presence of healthcare professionals and health care staff with the active participation of the research team is based on the initial data provided in the first phase of the study Phase IV: Using a performance-based incentive system to achieve goals based on initial data. In this regard, the focus group (FGD) method is used to reach the best option for an ideal incentive. In this phase only one group will arrive. * It should be noted that 4 rural centers and 4 urban centers will not receive any intervention.

Settings and conduct

This study will be carried out in Bushehr (Dashtestan), Semnan (Damqan and Garmsar) and Iran (shahryar) universities of medical sciences, .

Participants/Inclusion and exclusion criteria

16 urban health centers and 16 rural health homes

Intervention groups

16 urban health centers and 16 health homes

Main outcome variables

Hypertension; diabetes; insufficient physical activity; tobacco smoking; insufficient intake of fruits/vegetables; body mass index

General information

Reason for update

revised based on reviewer comment (of course, it was done in consultation with an epidemiologist)

Acronym

IRPONT

IRCT registration information

IRCT registration number: **IRCT20081205001488N2**

Registration date: **2018-06-03, 1397/03/13**

Registration timing: **prospective**

Last update: **2020-04-12, 1399/01/24**

Update count: **2**

Registration date

2018-06-03, 1397/03/13

Registrant information

Name

Maziar Moradi-Lakeh

Name of organization / entity

Iran University of Medical Sciences

Country

Iran (Islamic Republic of)

Phone

+98 21 8860 2225

Email address

moradilakeh.m@iums.ac.ir

Recruitment status

Recruitment complete

Funding source

National Institute for Medical Research Development (NIMAD)

Expected recruitment start date

2018-06-17, 1397/03/27

Expected recruitment end date

2021-03-18, 1399/12/28

Actual recruitment start date

empty

Actual recruitment end date

empty

Trial completion date

empty	we use randomization to randomly assign centers for intervention groups
Scientific title	Blinding (investigator's opinion)
The impact of a results-based motivating system on population levels of the non-communicable diseases risk factors in Iran: A field trial study	Not blinded
Public title	Blinding description
The impact of a results-based motivating system on population levels of the non-communicable diseases risk factors in Iran: A field trial study	Placebo
Purpose	Used
Health service research	Assignment
Inclusion/Exclusion criteria	Parallel
Inclusion criteria:	Other design features
In the first study, three medical universities will be selected randomly from three different climates, and they will be asked to submit a list of health and medical and health centers in their urban and rural areas, broken down by health centers and health homes. Then the list of networks that have the entry criteria (descriptions in the entry criteria) are prepared and then the selection is made randomly. From each University, four Urban health centers and four health home randomly selected . Four groups (each consisting of a health center and a health center) will be introduced.Of course, to control the likelihood of data contamination, one of the universities(Semnan), in addition to the intervention cluster city, will select another city(Garmsar) with 4 urban clusters and 4 rural clusters, which will receive no intervention. Universities eligible for entry into the study: 1. Have the consent to cooperate in the study. Cities eligible for entry into the study are: 1. Cities with at least four urban health centers and four health-care homes. Eligible entry centers: 1. Health homes with at least two "Behvarz". 2. City bases Have at least 2 health care staff. And preferably have recruiting staff . 3. health homes that are preferable to the Very small village .eligible staff : Preferably, have fixed forces (recruitment) of health centers and health homes At least 2 years into that center.	This is a field trial. Interventions are performed at the first health level of the selected areas, and the final outcome is measured using population surveys.
Exclusion criteria:	
Disagreement with the text of the memorandum, by the relevant authorities of the universities	
Age	Secondary Ids
From 30 years old to 70 years old	empty
Gender	Ethics committees
Both	1
Phase	Ethics committee
N/A	Name of ethics committee
Groups that have been masked	national institute for medical research development
No information	Street address
Sample size	Tehran, West Fatemi St., Besat Street, No. 21
Target sample size: 32	City
More than 1 sample in each individual	Tehran
Number of samples in each individual: 40	Province
Thirty-two clusters are considered and for each cluster, 40 samples (individuals with inclusion criteria) will be provided.	Tehran
Randomization (investigator's opinion)	Postal code
Randomized	66900920-66938037
Randomization description	Approval date
	2017-07-31, 1396/05/09
	Ethics committee reference number
	IR.NIMAD.REC.1396.084
	Health conditions studied
	1
	Description of health condition studied
	Risk factors of Non-communicable diseases
	ICD-10 code
	ICD-10 code description
	Primary outcomes
	1
	Description
	Population level of uncontrolled hypertension
	Timepoint
	At the beginning of intervention (Month 0), 12 months after starting of intervention, 24 month after starting of intervention
	Method of measurement
	Population Survey

2

Description

Population level of Poorly controlled diabetes

Timepoint

At the beginning of intervention (Month 0), 12 months after starting of intervention, 24 month after starting of intervention

Method of measurement

Population Survey

3

Description

Population level of insufficient physical activity

Timepoint

At the beginning of intervention (Month 0), 12 months after starting of intervention, 24 month after starting of intervention

Method of measurement

Population Survey

4

Description

Population level of current tobacco smoking

Timepoint

At the beginning of intervention (Month 0), 12 months after starting of intervention, 24 month after starting of intervention

Method of measurement

Population Survey

5

Description

Population level of insufficient intake of fruits/vegetables

Timepoint

At the beginning of intervention (Month 0), 12 months after starting of intervention, 24 month after starting of intervention

Method of measurement

Population Survey

6

Description

Population level of body mass index

Timepoint

At the beginning of intervention (Month 0), 12 months after starting of intervention, 24 month after starting of intervention

Method of measurement

Population Survey

Secondary outcomes

empty

Intervention groups

1

Description

Group IV: Assessment of the main NCDs' risk factors and setting time-bound targets, AND Finding and sharing evidence on effective/efficient interventions for controlling the risk factors AND Operational planning with contribution of local health authority

Category

Other

2

Description

Group I: Assessment of the main NCDs' risk factors and setting time-bound targets.

Category

Other

3

Description

Group II: Assessment of the main NCDs' risk factors and setting time-bound targets AND Finding and sharing evidence on effective/efficient interventions for controlling the risk factors

Category

Other

4

Description

Group III: Assessment of the main NCDs' risk factors and setting time-bound targets AND Finding and sharing evidence on effective/efficient interventions for controlling the risk factors AND Operational planning with contribution of local health authority

Category

Other

5

Description

Control group: A county (Garmsar County) consisting of 4 health houses and 4 health center will not receive any intervention, and only three survey will be conducted.

Category

Other

Recruitment centers

1

Recruitment center

Name of recruitment center

Iran university of medical sciences

Full name of responsible person

Maziar Moradi-Lakeh

Street address

Shahriar Health Network

City

Shahriar

Province

Tehran
Postal code
1449614535
Phone
+98 21 8860 2225
Email
mazmoradi@gmail.com

2

Recruitment center

Name of recruitment center
Bushehr University of Medical Sciences
Full name of responsible person
Maziar Moradi-Lakeh
Street address
Sports ST, Bushehr
City
Borazjan
Province
Boushehr
Postal code
1449614535
Phone
+98 71 3252 2078
Email
mazmoradi@gmail.com

3

Recruitment center

Name of recruitment center
Semnan University of Medical Sciences
Full name of responsible person
Maziar Moradi-Lakeh
Street address
Basij Blvd
City
Damghan
Province
Semnan
Postal code
3519899951
Phone
+98 23 3344 1022
Email
mazmoradi@gmail.com

4

Recruitment center

Name of recruitment center
Semnan University of Medical Sciences
Full name of responsible person
Maziar Moradi-Lakeh
Street address
Shahid Rajaie Street
City
Garmsar
Province
Semnan
Postal code
3519899951

Phone
+98 23 3422 8830
Email
mazmoradi@gmail.com

Sponsors / Funding sources

1

Sponsor

Name of organization / entity
National Institute for Medical Research Development
(NIMAD)
Full name of responsible person
Dr. Sayena Rafizadeh - Project #958058
Street address
No 21, Besat St, West Fatemi Ave
City
Tehran
Province
Tehran
Postal code
1449614535
Phone
+98 21 8860 2225
Email
Mazmoradi@gmail.com

Grant name

Grant code / Reference number

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

Yes

Title of funding source

National Institute for Medical Research Development
(NIMAD)

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

Other

Person responsible for general inquiries

Contact

Name of organization / entity
Preventive Medicine and Public Health Research
Center
Full name of responsible person
Maziar Moradi-Lakeh
Position
Professor
Latest degree
Specialist
Other areas of specialty/work
Public Health/Community Medicine
Street address

IUMS, Crossroads of Hemmat-Chamran expressways,
Tehran, Iran

City

Tehran

Province

Tehran

Postal code

1449614535

Phone

+98 21 8860 2225

Fax

Email

mazmoradi@yahoo.com

Web page address

Person responsible for scientific inquiries

Contact

Name of organization / entity

Preventive Medicine and Public Health Research
Center

Full name of responsible person

Dr. Maziar Moradi-Lakeh

Position

Professor

Latest degree

Specialist

Other areas of specialty/work

Public Health/Community Medicine

Street address

Iran University of Medical Sciences, Hemmat-
Chamran crossroads

City

Tehran

Province

Tehran

Postal code

1449614535

Phone

+98 21 8860 2225

Fax

Email

Mazmoradi@yahoo.com

Web page address

Person responsible for updating data

Contact

Name of organization / entity

Preventive Medicine and Public Health Research
Center

Full name of responsible person

Maziar Moradi-Lakeh

Position

Professor

Latest degree

Specialist

Other areas of specialty/work

Public Health/Community Medicine

Street address

IUMS, Crossroads of Hemmat-Chamran expressway

City

Tehran

Province

Tehran

Postal code

1635883813

Phone

+98 21 8860 2225

Fax

Email

mazmoradi@yahoo.com

Web page address

Sharing plan

Deidentified Individual Participant Data Set (IPD)

Yes - There is a plan to make this available

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

Yes - There is 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

Yes - There is a plan to make this available

Title and more details about the data/document

Access to relevant files is possible one year after the
publication, with correspondence email:
mazmoradi@gmail.com.

When the data will become available and for how long

One year after publication

To whom data/document is available

Academic researchers

Under which criteria data/document could be used

There is no limitation to the analysis

From where data/document is obtainable

Contact with email: mazmoradi@gmail.com

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

Study aims of the study, about 10 to 21 days.

Comments