

# Clinical Trial Protocol

## Iranian Registry of Clinical Trials

09 Jun 2026

### Design, Implementation and Evaluation of Preoperative Patient Self-Efficacy Training Program in Ostomy Care

#### Protocol summary

##### Study aim

Design, Implementation and Evaluation of Preoperative Patient Self-Efficacy Training Program in Ostomy Care

##### Design

Clinical trial with control group, with parallel groups, on 80 patients

##### Settings and conduct

80 patients who are in the colostomy or ileostomy surgery list in the centers affiliated to Shahid Beheshti University of Medical Sciences will be divided into two groups (intervention and control) by simple random allocation method. patients in the intervention group, In addition to training in the usual way, they will also be trained with the ostomt self-efficacy training program before surgery. the patients of the control group only receive education in the usual way, i.e. education after surgery. Demographic questionnaire tools, clinical information questionnaire and stoma self-efficacy scale will be completed by the patients before surgery and one month after surgery.

##### Participants/Inclusion and exclusion criteria

Inclusion: Age 18 to 65 years, Absence of cognitive or psychological impairment, Ability to communicate,

Exclusion: Performing emergency surgery

##### Intervention groups

Intervention group patients: In addition to training in the usual way, they are also trained with the self-efficacy training program of the ostomy prepared before the surgery. The training session is conducted face-to-face for each patient and, if the patient wishes, with the presence of his family or caregiver. The session will last for 45 minutes in a hospital or clinic by a wound and ostomy nurse within a week before surgery. Patients of the control group only receive education in the usual way, i.e. education after surgery.

##### Main outcome variables

Ostomy self-efficacy; Hospitalization time; Number of readmissions; Number of emergency visits; The peristomal skin condition

#### General information

##### Reason for update

##### Acronym

##### IRCT registration information

IRCT registration number: **IRCT20220731055588N1**

Registration date: **2022-08-09, 1401/05/18**

Registration timing: **registered\_while\_recruiting**

Last update: **2022-08-09, 1401/05/18**

Update count: **0**

##### Registration date

2022-08-09, 1401/05/18

##### Registrant information

##### Name

ali ali bakhoda

##### Name of organization / entity

##### Country

Iran (Islamic Republic of)

##### Phone

+98 21 8865 5379

##### Email address

ali.baakhoda@gmail.com

##### Recruitment status

##### Recruitment complete

##### Funding source

##### Expected recruitment start date

2022-08-06, 1401/05/15

##### Expected recruitment end date

2023-06-21, 1402/03/31

##### Actual recruitment start date

empty

##### Actual recruitment end date

empty

##### Trial completion date

empty

##### Scientific title

Design, Implementation and Evaluation of Preoperative Patient Self-Efficacy Training Program in Ostomy Care

#### Public title

Design, Implementation and Evaluation of Preoperative Patient Self-Efficacy Training Program in Ostomy Care

#### Purpose

Education/Guidance

#### Inclusion/Exclusion criteria

##### Inclusion criteria:

Age 18 to 65 years  
Lack of cognitive or mental impairment  
Ability to verbal communicate

##### Exclusion criteria:

Performing emergency surgery

#### Age

From **18 years** old to **65 years** old

#### Gender

Both

#### Phase

N/A

#### Groups that have been masked

*No information*

#### Sample size

Target sample size: **80**

#### Randomization (investigator's opinion)

Randomized

#### Randomization description

Random allocation will be done using a coin toss by an independent researcher. To prepare the randomized list, a coin will be tossed 80 times. If the coin lands on A, it means the intervention group, and if it lands on B, it means the control group. The randomization results will be recorded in a list from 1 to 80 respectively. In order to hide the results from the researcher's view, according to the list, each of the results will be recorded on a sheet and will be placed in a sealed envelope. The list number corresponding to that result will be written on each envelope. After each sample enters the research, the envelopes from 1 to 80 will be opened and the placement of the patient in the intervention or control group will be determined.

#### Blinding (investigator's opinion)

Not blinded

#### Blinding description

#### Placebo

Not used

#### Assignment

Parallel

#### Other design features

### Secondary Ids

empty

### Ethics committees

#### 1

##### Ethics committee

###### Name of ethics committee

Ethics Committee of Pharmacy, Nursing and Midwifery Faculties of Shahid Beheshti University of

Medic

#### Street address

School of Nursing and Midwifery, Shahid Beheshti University of Medical Sciences, No. 2660, Vali Asr Ave., Niayesh Cross Road, Tehran, Iran

#### City

Tehran

#### Province

Tehran

#### Postal code

1985717443

#### Approval date

2022-06-21, 1401/03/31

#### Ethics committee reference number

IR.SBMU.PHARMACY.REC.1401.060

### Health conditions studied

#### 1

##### Description of health condition studied

Colostomy; Ileostomy; Colon Malignancies

##### ICD-10 code

##### ICD-10 code description

### Primary outcomes

#### 1

##### Description

The ostomy self-efficacy: The Ostomy self-efficacy is equal to the score obtained by the patient in the ostomy self-efficacy questionnaire. The questionnaire has 22 items in a 5-item Likert scale. The questionnaire is in two dimensions includes ostomy care self-efficacy (13 items) and social self-efficacy (9 items). A higher score indicates higher self-efficacy. The lowest score that can be obtained in this tool is 22 and the highest score is 110.

##### Timepoint

The stoma self-efficacy questionnaire will complete by patients before surgery and one month after surgery.

##### Method of measurement

The stoma self-efficacy scale was designed and psychometrically evaluated by Beckers et al. (1996). The questionnaire has 22 items in two dimensions, including ostomy care self-efficacy (first 13 items) and social self-efficacy (next 9 items). Using a Likert scale, the answers are defined from not sure at all (score 1) to completely sure (score 5). A higher score indicates greater ostomy self-efficacy. The lowest score that can be obtained in this tool is 22 and the highest score is 110.

### Secondary outcomes

#### 1

##### Description

Hospitalization time

##### Timepoint

One month after surgery

##### Method of measurement

Clinical records and patient interview

## 2

### **Description**

The number of readmissions

### **Timepoint**

One month after surgery

### **Method of measurement**

Clinical records and patient interview

## 3

### **Description**

The number of emergency visits

### **Timepoint**

One month after surgery

### **Method of measurement**

Clinical records and patient interview

## 4

### **Description**

Peristomal skin condition

### **Timepoint**

One month after surgery

### **Method of measurement**

DET ostomy skin tool

## **Intervention groups**

### 1

#### **Description**

Intervention group: (Pre-surgery training In addition to usual training) The patients who are allocated in the intervention group will be invited to participate in an ostomy care training session within one week before surgery in the one of clinic visit sessions or during hospitalization. A training session for 45 minutes will be held in the hospital or clinic for each patient independently. The researcher along with a wound and ostomy nurse will lead the training session. The goals of the training session will include a) familiarizing the patient with the educational content at the level of cognitive perception and b) the ability to change the ostomy bag and take care of the skin around the ostomy at the level of imitation of the psychomotor domain. Educational content will be presented to patients by using lectures, practical simulation of ostomy bag replacement skills, presentation of educational booklets to patients and introduction of online pages in virtual social networks to learn more. Content will be presented in simple and understandable language for the patient. The educational booklet and online pages of social networks will be prepared by the researcher based on the educational content and with the final approval of the faculty members of Shahid Beheshti University of Medical Sciences. The online pages of social networks will contain uploaded texts, images and educational videos; The membership link will only be available to patients of the intervention group or one of their family members (if the patient wishes). The time of use of

educational materials will be at the disposal of the patient, who can adjust it according to his needs. The ostomy bag replacement simulation tool will include one-piece and two-piece ostomy bag, ostomy size measurement template, scissors, marker, gloves, ostomy powder, ostomy paste, ostomy skin protection ring, ostomy belt, intestinal ostomy simulator. The patient will have Practical work on the simulator under the guidance of a wound and ostomy nurse to familiarize with how to change the ostomy bag and related tips to prevent skin complications related to ostomy. The patient's family and their caregivers will be allowed to attend the training session.

#### **Category**

Other

## 2

#### **Description**

Control group: (Usual training) For the patients in the control group, there will be no training by the research team. These patients will be trained in the usual way, i.e. after surgery and before discharge.

#### **Category**

Other

## **Recruitment centers**

### 1

#### **Recruitment center**

##### **Name of recruitment center**

Taleghani Hospital (Tehran)

##### **Full name of responsible person**

Dr. Hasani

##### **Street address**

A-a-rabi Ave.

##### **City**

Tehran

##### **Province**

Tehran

##### **Postal code**

1985711151

##### **Phone**

+98 21 2243 2560

##### **Email**

taleghanihospital@sbmu.ac.ir

### 2

#### **Recruitment center**

##### **Name of recruitment center**

Imam Hosein Hospital (Tehran)

##### **Full name of responsible person**

Dr. Sadeghi

##### **Street address**

Shaheed Madani Ave.

##### **City**

Tehran

##### **Province**

Tehran

##### **Postal code**

1617763141

**Phone**

+98 21 7755 8001

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info@ehmc.ir

**3**

**Recruitment center**

**Name of recruitment center**

Shohadaye Tajrish Hospiat (Tehran)

**Full name of responsible person**

Dr. Abdollahi Majd

**Street address**

Shahrdari Ave.

**City**

Tehran

**Province**

Tehran

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1989934148

**Phone**

+98 21 25719

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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**

Vice President of Research and Technology

**Street address**

School of Nursing and Midwifery, Shahid Beheshti University of Medical Sciences, No. 2660, Vali Asr Ave., Niayesh Cross Road, Tehran, Iran

**City**

Tehran

**Province**

Tehran

**Postal code**

1985717443

**Phone**

+98 21 8865 5366

**Email**

sbnmf@sbmu.ac.ir

**Web page address**

<https://nm.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

**Person responsible for general inquiries**

**Contact**

**Name of organization / entity**

Shahid Beheshti University of Medical Sciences

**Full name of responsible person**

Mahnaz Ilkhani

**Position**

Associate Professor

**Latest degree**

Ph.D.

**Other areas of specialty/work**

Nursing

**Street address**

School of Nursing and Midwifery, Shahid Beheshti University of Medical Sciences, No. 2660, Vali Asr Ave., Niayesh Cross Road, Tehran, Iran

**City**

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**Province**

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**Postal code**

1985717443

**Phone**

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**Email**

m\_ilkhani@yahoo.com

**Person responsible for scientific inquiries**

**Contact**

**Name of organization / entity**

Shahid Beheshti University of Medical Sciences

**Full name of responsible person**

Mahnaz Ilkhani

**Position**

Associate Professor

**Latest degree**

Ph.D.

**Other areas of specialty/work**

Nursing

**Street address**

School of Nursing and Midwifery, Shahid Beheshti University of Medical Sciences, No. 2660, Vali Asr Ave., Niayesh Cross Road, Tehran, Iran

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**Province**

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**Postal code**

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**Phone**

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**Email**

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

Shahid Beheshti University of Medical Sciences

**Full name of responsible person**

Mahnaz Ilkhani

**Position**

Associate Professor

**Latest degree**

Ph.D.

**Other areas of specialty/work**

Nursing

**Street address**

School of Nursing and Midwifery, Shahid Beheshti  
University of Medical Sciences, No. 2660, Vali Asr  
Ave., Niayesh Cross Road, Tehran, Iran

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**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**

All data collected from assessed variables can be shared  
after de-identification.

**When the data will become available and for how long**

As soon as the article is published, it can be shared.

**To whom data/document is available**

All researchers and interested parties are allowed to  
receive research data.

**Under which criteria data/document could be used**

Sending a request via email is required to receive data  
by researchers and interested parties.

**From where data/document is obtainable**

Sending a request via email is required to receive data  
by researchers and interested parties. email:  
m\_ilkhani@yahoo.com

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

Sending a request via email is required to receive data  
by researchers and interested parties.

**Comments**