

Clinical Trial Protocol

Iranian Registry of Clinical Trials

13 Jul 2026

Comparing the effectiveness of integrated care versus usual care on maternal grief in mothers with perinatal loss: a randomized clinical trial

Protocol summary

Study aim

Evaluation of the effect of integrated care compared to usual care on maternal greif in bereaved mothers with perinatal loss

Design

A single-blind, parallel-group clinical trial with a control group, randomized by a randomization computer program with 128 participants, 64 in each group.

Settings and conduct

The setting of the study will be the maternity ward of Bahar Hospital in Shahroud city of Semnan province. The study population is hospitalized mothers with perinatal fetal loss. Blinding will be done centrally by a statistician and the list of blocks will be made available to someone outside the research. This person will prepare envelopes containing the name of intervention or control in the order of the list and number them in order. Then the envelopes will be given to the person in charge of the delivery room. After obtaining informed consent from the bereaved mothers, the first thick sealed envelope will be given to the principal researcher by the delivery room manager from the numbered box.

Participants/Inclusion and exclusion criteria

Inclusion criterion: mothers with pregnancy loss.

Exclusion criteria: Voluntary or legal abortion of pregnancy with clearly abnormal fetus, Having a mental illness or a history of it

Intervention groups

The intervention group: they will have a face-to-face consultation at the time of discharge from the hospital, and a virtual group counseling session will be conducted 20 to 35 days after discharge. From the time of discharge, they have been a member of the virtual information network for 2 months and will be able to receive training and advice. Control group: At the time of discharge, they are not aware of their group and will receive only a pamphlet.

Main outcome variables

Maternal grief, psychological well-being

General information

Reason for update

Acronym

IRCT registration information

IRCT registration number: **IRCT20230907059370N1**

Registration date: **2023-10-10, 1402/07/18**

Registration timing: **registered_while_recruiting**

Last update: **2023-10-10, 1402/07/18**

Update count: **0**

Registration date

2023-10-10, 1402/07/18

Registrant information

Name

giti Atashsokhan

Name of organization / entity

Country

Iran (Islamic Republic of)

Phone

+98 23 3239 5054

Email address

gitiatashsokhan@gmail.com

Recruitment status

Recruitment complete

Funding source

Expected recruitment start date

2023-09-29, 1402/07/07

Expected recruitment end date

2024-02-09, 1402/11/20

Actual recruitment start date

empty

Actual recruitment end date

empty

Trial completion date

empty

Scientific title

Comparing the effectiveness of integrated care versus usual care on maternal grief in mothers with perinatal loss: a randomized clinical trial	
Public title	Integrated care effect on grief of bereaved mothers
Purpose	Supportive
Inclusion/Exclusion criteria	
Inclusion criteria:	Mothers with perinatal loss Stillbirth Ectopic pregnancy Fetal death immediately after birth during the mother's hospitalization Abortion, miscarriage Having smart phone Being married
Exclusion criteria:	Suffering from mental illness Having a history of psychological illness Taking psychoactive drugs Divorce Experience a new crisis during past 6 months Pregnancy loss with abnormal fetus
Age	No age limit
Gender	Female
Phase	N/A
Groups that have been masked	• Participant • Care provider • Outcome assessor • Data analyser
Sample size	Target sample size: 128
Randomization (investigator's opinion)	Randomized
Randomization description	The random allocation of mothers is done using blocks of size four and by a Statistics specialist through computer-based randomization. Based on the table obtained from the allocation, envelopes that are numbered in sequence and contain the type of intervention or control are prepared by someone outside the research. The thick envelopes in the package are placed in a box in order. This box is given to the person in charge of the delivery room who is not involved in the research. After obtaining the consent of the participant to enter the study, an envelope will be opened by the delivery room manager. The desired sample will be assigned to the specified group inside the envelope. Participants will not know about the allocation
Blinding (investigator's opinion)	Single blinded
Blinding description	Participants (bereaved mothers) will receive the usual care, but the intervention group also will receive additional care. The intervention group will have face-to-face counseling and will have access to the virtual educational counseling network. The control group will only receive an educational pamphlet at the time of discharge from the hospital. Both groups are unaware of the allocation. Clinical caregivers will perform all routine tasks for both groups and will not be informed. The person responsible for data collection is a research assistant who will complete the questionnaires over the phone for both groups and will not be informed. The statistician will analyze the data from the questionnaires and will not be informed.
Placebo	Not used
Assignment	Parallel
Other design features	
Secondary Ids	empty
Ethics committees	
1	
Ethics committee	
Name of ethics committee	Ethics committee of Shahroud University of Medical Sciences
Street address	Shahroud University of Medical Sciences and Health Services, Hafte Tir Square, Taheri Ave
City	Shahroud
Province	Semnan
Postal code	3614773955
Approval date	2021-03-08, 1399/12/18
Ethics committee reference number	IR.SHMU.REC.1400.011
Health conditions studied	
1	
Description of health condition studied	Maternal grief, psychological well-being
ICD-10 code	
ICD-10 code description	
Primary outcomes	
1	
Description	Maternal greif score in Perinatal Grief Scale (potvin)
Timepoint	Before intervention, after virtual session, 2 months after discharge
Method of measurement	Perinatal Grief Scale
2	
Description	Psychological well-being

Timepoint

Before intervention, 2 months after discharge

Method of measurement

Ryff scales of psychological well-being

Secondary outcomes

empty

Intervention groups**1****Description**

Intervention group: one face-to-face individual counseling session patient-centered for bereaved mother with an empathic approach before discharge from the hospital, two virtual peer group counseling sessions, and allocation of a virtual online counseling channel for bereaved mothers until the end of the sampling period.

Category

Rehabilitation

2**Description**

Control group: receiving usual care and a general educational pamphlet at time of discharge

Category

Rehabilitation

Recruitment centers**1****Recruitment center****Name of recruitment center**

Maternity ward at Bahar hospital in Shahroud

Full name of responsible person

Giti Atashsokhan

Street address

Bahar Hospital, 22 Bahman street

City

Shahroud

Province

Semnan

Postal code

3614773955

Phone

+98 23 3222 7607

Fax**Email**

atashsokhan@shmu.ac.ir

Web page address

https://shmu.ac.ir/bahar/fa

Sponsors / Funding sources**1****Sponsor**

Name of organization / entity

Shahroud University of Medical Sciences

Full name of responsible person

Dr.Mohammad Hasan Imamian

Street address

7th Tir Square, Taheri St. Shahrood University of Medical Sciences.Shahroud, Iran.

City

Shahroud

Province

Semnan

Postal code

۳۶۱۴۷۷۳۹۴۳

Phone

+98 23 3239 4852

Fax**Email**

vcr@shmu.ac.ir

Web page address

https://shmu.ac.ir/research/fa

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

Yes

Title of funding source

Shahroud University of Medical Sciences

Proportion provided by this source

2

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**

Shahroud University of Medical Sciences

Full name of responsible person

Giti Atashsokhan

Position

faculty member

Latest degree

Master

Other areas of specialty/work

Reproductive Health

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7th Tir square, Taheri street

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Email

Person responsible for scientific inquiries

Contact

Name of organization / entity
Shahroud University of Medical Sciences

Full name of responsible person
Afsaneh Keramat

Position
Professor

Latest degree
Ph.D.

Other areas of specialty/work
Reproductive Health

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Person responsible for updating data

Contact

Name of organization / entity
Shahroud University of Medical Sciences

Full name of responsible person
Giti Atashsokhan

Position
Faculty member

Latest degree
Master

Other areas of specialty/work
Reproductive Health

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Phone
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Email
atashsokhan@shmu.ac.ir

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

Undecided - It is not yet known if there will be 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

Undecided - It is not yet known if there will be 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

After publishing the article and anonymizing the participants, the data file can be sent via email as requested

When the data will become available and for how long

After publishing the research paper of the project for 6 months

To whom data/document is available

for all Academic Researcher

Under which criteria data/document could be used

An approved proposal whose goals indicate the need for this information

From where data/document is obtainable

By Email

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

After the approval of the project partners, the information will be sent within two weeks

Comments