

Clinical Trial Protocol

Iranian Registry of Clinical Trials

09 Jul 2026

To evaluate Neoadjuvant chemotherapy followed by chemoradiation in patients with unresectable cervical cancer

Protocol summary

Registration timing: **prospective**

Study aim

To evaluate response to treatment of neoadjuvant chemotherapy followed by chemoradiation in patients with unresectable cervical cancer

Last update: **2021-10-21, 1400/07/29**

Update count: **0**

Registration date

2021-10-21, 1400/07/29

Design

Phase II clinical trial with single group including 74 subjects who referred and follow up for 3 months after treatment

Registrant information

Name

Marziye Heidarpourfard

Name of organization / entity

Country

Iran (Islamic Republic of)

Phone

+98 71 3823 5257

Email address

heidarpourfard@sums.ac.ir

Settings and conduct

Newly diagnosed unresectable cervical cancer patients who referred to Shiraz radiation oncology department received four cycle chemotherapy in form of carboplatin with (Area under curve AUC:5) and paclitaxel(175 per square meter) every 3 weeks after obtaining informed consent, then chemoradiation with External beam radiotherapy and brachytherapy up to 80-90 Gy to high risk clinical target volume and concurrent chemotherapy with cisplatin (40 mg per square meter) and then response to treatment evaluate three months after treatment by recist1.1 guideline with MRI

Recruitment status

Recruitment complete

Funding source

Participants/Inclusion and exclusion criteria

Newly diagnosed unresectable cervical cancer without history of metastasis and adequate bone marrow, hepatic, renal function and age 18 to 75 years and performance status 0 and 1 without Previous pelvic radiation, Serious medical illness, hypersensitivity to chemotherapy agent

Expected recruitment start date

2021-10-23, 1400/08/01

Expected recruitment end date

2022-06-21, 1401/03/31

Actual recruitment start date

empty

Actual recruitment end date

empty

Trial completion date

empty

Intervention groups

single arm group treated by neoadjuvant chemotherapy

Scientific title

To evaluate Neoadjuvant chemotherapy followed by chemoradiation in patients with unresectable cervical cancer

Main outcome variables

Response to treatment

Public title

To evaluate Neoadjuvant chemotherapy in treatment of unresectable cervical cancer patients

General information

Reason for update

Acronym

IRCT registration information

IRCT registration number: **IRCT20210808052110N1**

Registration date: **2021-10-21, 1400/07/29**

Purpose

Treatment

Inclusion/Exclusion criteria

Inclusion criteria:

1.Unresectable cervical cancer(IIB toIVA,IB3,IIA more than 4cm) 2.No history of metastasis 3.Adequate renal function(GFR more than 60ml per min per1.73) 4.Adequate hepatic function(Liver enzyme<5* upper limit normal) 5.Adequate bone marrow function(Absolute neutrophil>1500) 6-Without history of systemic disease 7.Absence of active infection 8.Absence of previous or concurrent oncology pathology 9.ECOG performance status 0 and 1 10.Age 18 to 75 years

Exclusion criteria:

1-Previous pelvic radiation 2-Pregnancy 3-Serious medical illness 4-Hypersensitivity to chemotherapy agent

Age

From **18 years** old to **75 years** old

Gender

Female

Phase

2

Groups that have been masked

No information

Sample size

Target sample size: **74**

Randomization (investigator's opinion)

N/A

Randomization description

Blinding (investigator's opinion)

Not blinded

Blinding description

Placebo

Not used

Assignment

Single

Other design features

Secondary Ids

empty

Ethics committees

1

Ethics committee

Name of ethics committee

Ethics committee of Shiraz university of medicine

Street address

Shiraz university of medicine-in front of Felestin street-Zand street-Shiraz

City

Shiraz

Province

Fars

Postal code

7134814336

Approval date

2021-06-21, 1400/03/31

Ethics committee reference number

IR.SUMS.MED.REC.1400.173

Health conditions studied

1

Description of health condition studied

cancer of cervix

ICD-10 code

C53

ICD-10 code description

Malignant neoplasm of cervix uteri

Primary outcomes

1

Description

Response to treatment 3 months after fulfilling it

Timepoint

3 months after fulfilling treatment

Method of measurement

To do MRI and evaluate response to treatment by recist 1.1 guideline

Secondary outcomes

empty

Intervention groups

1

Description

Intervention group:Receiving 4 course chemotherapy with carboplatin (Area under curve ;AUC:5)and paclitaxel (175mg per square meter)every three week before chemoradiation with External beam radiation and brachytherapy up to 80 to 90Gy to high risk Clinical target volume (CTV)with prescribing concurrent cisplatin weekly(40mg per square meter)

Category

Treatment - Drugs

Recruitment centers

1

Recruitment center

Name of recruitment center

Radiation oncology department of Shiraz university of medicine

Full name of responsible person

Marzieh Heidarpourfard

Street address

Namazi hospital-Namazi square-Zand street-Shiraz

City

Shiraz

Province

Fars

Postal code

7193613311

Phone

+98 71 3647 4332

Fax
+98 71 3647 4326
Email
nemazee_inf@sums.ac.ir

Sponsors / Funding sources

1

Sponsor

Name of organization / entity
Shiraz University of Medical Sciences
Full name of responsible person
Yones Ghasemi
Street address
7th floor-Shiraz university of medical science-Zand
street
City
Shiraz
Province
Fars
Postal code
7134814336
Phone
+98 71 3230 5410
Fax
Email
vcrdep@sums.ac.ir

Grant name

Grant code / Reference number

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

Yes

Title of funding source

Shiraz 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
Shiraz University of Medical Sciences
Full name of responsible person
Marzieh Heidarpourfard
Position
Resident
Latest degree
Medical doctor
Other areas of specialty/work
Radiotherapy
Street address
4th floor-Yas building-15/2 alley-Namazi street-

Sattarkan Blvd

City

Shiraz

Province

Fars

Postal code

7184763955

Phone

+98 71 3648 8332

Email

heidarpourfard@sums.ac.ir

Person responsible for scientific inquiries

Contact

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Shiraz University of Medical Sciences
Full name of responsible person
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City
Shiraz
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Fars
Postal code
7184763955
Phone
+98 71 3648 8332
Email
heidarpourfard@sums.ac.ir

Person responsible for updating data

Contact

Name of organization / entity
Shiraz University of Medical Sciences
Full name of responsible person
Marzieh Heidarpourfard
Position
Resident
Latest degree
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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

Not applicable

Analytic Code

Not applicable

Data Dictionary

Yes - There is a plan to make this available

Title and more details about the data/document

Individual data of study participants and all data can be shared after unrecognized them

When the data will become available and for how long

Access period starts 12 months after the results are published

To whom data/document is available

Researchers of university scientific institutes

Under which criteria data/document could be used

Any kind of analysis can be done on the delivered data after it is clarified and proved

From where data/document is obtainable

Marzie Heidarpourfard, email
address:heidarpourfard@sums.ac.ir

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

Please send me an email explaining your request for which part of the data you need and description of the analysis in question. If I see it is fit, the data will be emailed to you within a week

Comments