

Clinical Trial Protocol

Iranian Registry of Clinical Trials

23 Feb 2026

Comparison of the therapeutic effect of Sofosbuvir with Control group in outpatients with Covid-19 referred to outpatient clinics in Ahvaz

Protocol summary

Clinical recovery (composite) within 14 days

Study aim

To evaluate whether (Sofosbuvir) increases significant clinical improvement as compared to standard of care in out patients patients with mild COVID-19

Design

This is a parallel 2-arm randomized, controlled, double-blind, multi center study. 360 patients are enrolled and followed for 14 days

Settings and conduct

This study will be performed in outpatient medical centers in Ahvaz by researchers from the Lung Diseases Research Institute. Physicians who evaluate results and statistics that analyze data become blind, and patients and physicians who treat patients do not know the specific treatment group.

Participants/Inclusion and exclusion criteria

Inclusion criteria: Outpatients with positive PCR and mild clinical signs without pulmonary involvement on CT scan. (As defined by patient with sore throat, dry cough \pm , loss of sense of smell and taste, body aches, nausea and vomiting, O₂ Sat \geq 95, T $<$ 38, RR $<$ 24, stable pulse and blood pressure). Exclusion criteria: 18y $>$ age, transplant patients, severe skin history including TEN or Steven Johnson, allergy or allergy to the drug Male study, the patient has previously been treated with this drug for other reasons (hepatitis C, etc.) , Is not able to take pills, is not aware of his condition, is deficient in G6PD enzyme, is being treated with anticonvulsant drugs, has fever and respiratory symptoms for 2 weeks, is being treated with immunosuppressive drugs, 1 month ago drugs Has received antiviral for Covid-19, is pregnant or lactating, has ESRD, Advance CKD (liver, lung, heart, hematology) and solid tumors and cancers of neurological diseases such as CVA, etc., are excluded.

Intervention groups

360ligible patients with mild COVID-19 in a 1:1 ratio: • Standard of care treatment • tablet (Sofosbuvir 400mg) + Standard of care treatment

Main outcome variables

General information

Reason for update

Acronym

IRCT registration information

IRCT registration number: **IRCT20201127049505N1**

Registration date: **2021-02-06, 1399/11/18**

Registration timing: **retrospective**

Last update: **2021-02-06, 1399/11/18**

Update count: **0**

Registration date

2021-02-06, 1399/11/18

Registrant information

Name

mahtab moinpoor

Name of organization / entity

Country

Iran (Islamic Republic of)

Phone

+98 61 3322 1251

Email address

mahtab_moinpoor@yahoo.com

Recruitment status

Recruitment complete

Funding source

Expected recruitment start date

2020-12-28, 1399/10/08

Expected recruitment end date

2021-01-11, 1399/10/22

Actual recruitment start date

empty

Actual recruitment end date

empty

Trial completion date

empty

Scientific title

Comparison of the therapeutic effect of Sofosbuvir with Control group in outpatients with Covid-19 referred to outpatient clinics in Ahvaz

Public title

Study on the evaluation of the safety and efficacy of Sofosbuvir in participants with mild coronavirus in outpatients (COVID-19)

Purpose

Treatment

Inclusion/Exclusion criteria

Inclusion criteria:

Outpatients presented with positive PCR and mild clinical signs without pulmonary involvement on CT scan As defined by patient with sore throat dry cough loss of sense of smell and taste nausea and vomiting O₂ Sat \geq 95 T <38 RR <24 stable pulse and blood pressure body aches

Exclusion criteria:

y>18 Organ transplant Severe skin history including TEN or Steven Johnson Allergy to the studied drug The patient has previously been treated with this drug for other reasons Hepatitis C Not able to take pills Lack of awareness Be treated with anticonvulsant drugs Have fever and respiratory symptoms from 2 weeks ago Be treated with immunosuppressive drugs Be pregnant or breastfeeding 1 month ago received antiviral drugs for Covid-19 ESRD Advance CKD Advance liver disease Pulmonary Advance Disease Cardiac Advance Disease Advance Disease Hematology Solid tumor Cancer of neurological diseases such as CVA your situation G6PD deficient Be pregnant or breastfeeding

Age

From **18 years** old

Gender

Both

Phase

3

Groups that have been masked

No information

Sample size

Target sample size: **360**

Randomization (investigator's opinion)

Randomized

Randomization description

The nature of assigning patients to each of the two study groups was a randomized block method in which a 6-item block was used according to 60 6-item blocks. A: S B: c . In this way, patients are not aware of the treatment assigned to them. Also, the person evaluating the outcome of the treatment is kept unaware of the type of treatment. Blinding then Allocate Allocation In order to minimize the bias in the study, the Allocate method will be used. This will be done using unique codes or Unique code. In such a way that each person is assigned a three-digit number at random.

Blinding (investigator's opinion)

Single blinded

Blinding description

The task of treatment will remain unknown until the patient is random. Physicians who treat patients will remain blind. Physicians who evaluate results and statistics that analyze data will all become blind.

Placebo

Not used

Assignment

Parallel

Other design features

Secondary Ids

empty

Ethics committees

1

Ethics committee

Name of ethics committee

Ethics Committee of Ahwaz University of Medical Sciences

Street address

Imam Khomeini Hospital, 24 meters from the first floor of the office of the Department of Internal Education

City

ahwaz

Province

Khouzestan

Postal code

6193673116

Approval date

2020-12-23, 1399/10/03

Ethics committee reference number

IR.AJUMS.REC.1399.754

Health conditions studied

1

Description of health condition studied

COVID-19

ICD-10 code

U07.1

ICD-10 code description

COVID-19

Primary outcomes

1

Description

Shortness of breath

Timepoint

day 14 or sooner at the discretion of the physician

Method of measurement

Clinical evaluation

Secondary outcomes

empty

Intervention groups

1

Description

Intervention group:400mg of Sofosbovir and Vitamin B1300 control group

Category

Treatment - Drugs

2

Description

Control group: Vitamin B1300

Category

Placebo

Recruitment centers

1

Recruitment center

Name of recruitment center

Shafa Hospital

Full name of responsible person

Maryam Haddadzadeh Shoushtari

Street address

Imam Khomeini Hospital, 24 meters from the first floor of the office of the Department of Internal Education

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Ahwaz

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Khuzestan

Postal code

6193673116

Phone

+98 61 3322 1251

Email

drmahtab.moeinpour@yahoo.com

Sponsors / Funding sources

1

Sponsor

Name of organization / entity

Ahvaz University of Medical Sciences

Full name of responsible person

Mohammad Badavi Zade

Street address

Imam Khomeini Hospital, 24 meters from the first floor of the office of the Department of Internal Education

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Grant name

Grant code / Reference number

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

No

Title of funding source

not has

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

Ahvaz University of Medical Sciences

Full name of responsible person

Maryam Hadad Zade Shoushtari

Position

Consultant

Latest degree

Specialist

Other areas of specialty/work

Internal Medicine

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Person responsible for updating data

Contact

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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

The individual data of the participants in the study of all potential data can be shared after the identifiable individuals

When the data will become available and for how long

Access period starts from 1400

To whom data/document is available

The data will be available only to researchers working in academic and scientific institutions

Under which criteria data/document could be used

Only SPSS data will be available.

From where data/document is obtainable

09133583977

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

not

Comments