

Clinical Trial Protocol

Iranian Registry of Clinical Trials

23 Feb 2026

Effectiveness evaluation of Tofacitinib plus Remdesivir in comparison with Remdesivir in the treatment of adult patients with severe COVID-19 A randomized double-blind placebo-included clinical trial

Protocol summary

Study aim

Study of Tofacitinib effectiveness in the treatment of severe COVID-19 patients

Design

Double-blind, randomized clinical trial with parallel control group, phase 2-3 on 60 patients. Sealed envelope web-page was used for randomization.

Settings and conduct

The study will be performed in the ICU of Razi Hospital in Rasht from March to June 2021. Sealed envelope web-page is used to allocate treatments to the two arms A and B. 60 patients will be placed in 15 blocks of 4. In intervention group (A) they will receive Tofacitinib 10 mg/day orally for 14 days + Remdesivir 100 mg/day intravenously and in control group (B) they will receive one placebo tablet/day orally for 14 days + Remdesivir 100 mg/day intravenously. Blinding type: Double-blinded. For this purpose an Intensive Care Fellowship prescribes the assigned codes to each patient.

Participants/Inclusion and exclusion criteria

Inclusion criteria: All patients with severe Covid-19 admitted to the ICU of Razi Hospital in Rasht from March to June 2021. Exclusion criteria: Lack of informed consent; Lack of patient cooperation; Having active tuberculosis, hepatitis B or hepatitis C

Intervention groups

Intervention group: Tofacitinib oral tablet 10 mg daily + Remdesivir IV 100 mg daily for 14 days or hospital discharge (whichever comes first) Control group: Placebo oral tablet daily + Remdesivir IV 100 mg daily for 14 days or hospital discharge (whichever comes first)

Main outcome variables

The time required to improve clinical symptoms and paraclinical measures within 14 days of starting treatment

General information

Reason for update

Acronym

IRCT registration information

IRCT registration number: **IRCT20200329046892N2**

Registration date: **2021-03-09, 1399/12/19**

Registration timing: **prospective**

Last update: **2021-03-09, 1399/12/19**

Update count: **0**

Registration date

2021-03-09, 1399/12/19

Registrant information

Name

Nematollah Ahangar

Name of organization / entity

Country

Iran (Islamic Republic of)

Phone

+98 13 3369 0099

Email address

n.ahangar@gums.ac.ir

Recruitment status

Recruitment complete

Funding source

Expected recruitment start date

2021-03-15, 1399/12/25

Expected recruitment end date

2021-06-15, 1400/03/25

Actual recruitment start date

empty

Actual recruitment end date

empty

Trial completion date

empty

Scientific title

Effectiveness evaluation of Tofacitinib plus Remdesivir in comparison with Remdesivir in the treatment of adult patients with severe COVID-19 A randomized double-blind placebo-included clinical trial

Public title

Effect of tofacitinib in COVID-19

Purpose

Treatment

Inclusion/Exclusion criteria**Inclusion criteria:**

All patients with severe Covid-19 admitted to the ICU of Razi Hospital in Rasht for period of March- June 2021 Age greater than or equal to 18 years Patients admitted with the following criteria: fever (oral temperature greater than 37.2 ° C), dry cough, severe tiredness or dyspnea At least one of the following criteria : positive PCR OR lung involvement on chest X-ray / CT scan Absolute lymphocyte count greater than or equal to 200 per cubic millimeter Absolute neutrophil count greater than or equal to 1000 per milliliter Cubic meters Hemoglobin above 8 grams per deciliter

Exclusion criteria:

Lack of informed consent Lack of patient cooperation Having pulmonary embolism or intravascular thrombosis Having another active infection Having diverticulitis and gastric ulcer Having tuberculosis or hepatitis B or active hepatitis C or HIV Any major drug interaction between routine patient's drugs with any of the study drugs Pregnancy and lactation Simultaneous presence in other research study

Age

From **18 years** old

Gender

Both

Phase

2-3

Groups that have been masked

- Participant
- Care provider
- Outcome assessor
- Data analyser

Sample size

Target sample size: **60**

Randomization (investigator's opinion)

Randomized

Randomization description

It is a block randomization type and the block with size of 4. Randomization tool: sealedenvelope.com. 60 patients will be assigned in 15 blocks of 4. In intervention group (A) they will receive Tofacitinib 10 mg/day orally for 14 days + Remdesivir 100 mg/day intravenously and in control group (B) they will receive one placebo tablet/day orally for 14 days + Remdesivir 100 mg/day intravenously. Concealment is done using the sealed envelope method.

Blinding (investigator's opinion)

Double blinded

Blinding description

In this study, participants (patients) and medical staff are

kept blind to the specificity of study groups (Double-blinded). For this purpose, an independent person from the research team and an Intensive Care Fellowship prescribes the assigned codes to each patient. To ensure the similarity between the drug and the placebo, the placebo is made by the same company. Study drugs will be placed in similar packages, and patients will receive pre-arranged interventions in the order in which they enter the study.

Placebo

Used

Assignment

Parallel

Other design features**Secondary Ids**

empty

Ethics committees**1****Ethics committee****Name of ethics committee**

Ethics committee of Guilan University of Medical Sciences

Street address

School of Medicine, Guilan University Complex, 7th Km Tehran Road

City

Rasht

Province

Guilan

Postal code

4199613769

Approval date

2021-03-04, 1399/12/14

Ethics committee reference number

IR.GUMS.REC.1399.630

Health conditions studied**1****Description of health condition studied**

COVID-19

ICD-10 code

U07.1

ICD-10 code description

virus identified

2**Description of health condition studied**

COVID-19

ICD-10 code

U07.2

ICD-10 code description

virus not identified; Clinically-epidemiologically diagnosed COVID-19; Probable COVID-19; Suspected COVID-19

Primary outcomes

1

Description

The time required to improve clinical symptoms and within 14 days of treatment start

Timepoint

Daily from the first day of intervention

Method of measurement

Physical examination

Secondary outcomes

1

Description

Body temperature

Timepoint

Daily from the first day of intervention

Method of measurement

Thermometer

2

Description

Heart rate

Timepoint

Daily from the first day of intervention

Method of measurement

Pulse oxymeter

3

Description

Pulmonary radiological state

Timepoint

Weekly

Method of measurement

CT scan

4

Description

Blood pressure

Timepoint

Daily from the first day of intervention

Method of measurement

Barometer

5

Description

Mechanical ventilation

Timepoint

Daily from the first day of intervention

Method of measurement

Physical examination

6

Description

Duration of hospitalization in the intensive care unit

Timepoint

Daily from the first day of intervention

Method of measurement

Record in the patient file

7

Description

Mortality

Timepoint

Daily from the first day of intervention

Method of measurement

Record in the patient file

8

Description

Respiration rate

Timepoint

Daily from the first day of intervention

Method of measurement

Count

9

Description

Intubation

Timepoint

Daily from the first day of intervention

Method of measurement

Physical examination

10

Description

SPO2

Timepoint

Daily from the first day of intervention

Method of measurement

Pulse-oxy meter

11

Description

D-Dimer

Timepoint

Every 3 days from the first day of intervention

Method of measurement

Fluorescence immunochromatography

12

Description

Creatine phosphokinase

Timepoint

Every 3 days from the first day of intervention

Method of measurement

International Federation of Clinical Chemistry (IFCC)

13

Description

C reactive protein

Timepoint

Every 3 days from the first day of intervention

Method of measurement

Turbidometry

14

Description

Erythrocyte sedimentation rate

Timepoint

Every 3 days from the first day of intervention

Method of measurement

Westergren method

15

Description

Ferritin

Timepoint

Every 3 days from the first day of intervention

Method of measurement

Chemiluminescence method

Intervention groups

1

Description

Intervention group: Tofacitinib10 mg (Rofanib, Nano Alvand) 1 oral tablet daily + Remdesivir IV 100 mg daily for 14 days or hospital discharge (whichever comes first)

Category

Treatment - Drugs

2

Description

Control group: Placebo (Nano Alvand daru) 1 oral tablet daily + Remdesivir IV 100 mg daily for 14 days or hospital discharge (whichever comes first)

Category

Placebo

Recruitment centers

1

Recruitment center

Name of recruitment center

Razi hospital

Full name of responsible person

Mohammad Haghghi

Street address

Sardar Jangal Ave.

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Guilan

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4199613769

Phone

+98 13 3354 1001

Email

manesthesist@gmail.com

Sponsors / Funding sources

1

Sponsor

Name of organization / entity

Rasht University of Medical Sciences

Full name of responsible person

Dr. Mohammadreza Naghipour

Street address

Shahid Siadati Ave.. Namju St.

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Rasht

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4144666949

Phone

+98 13 3333 6394

Email

research@gums.ac.ir

Web page address

<https://www.gums.ac.ir/research>

Grant name

Grant code / Reference number

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

Yes

Title of funding source

Rasht 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

Rasht University of Medical Sciences

Full name of responsible person

Nematollah Ahangar

Position

Associate Professor

Latest degree

Ph.D.

Other areas of specialty/work

Pharmacology

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Person responsible for scientific inquiries**Contact****Name of organization / entity**

Rasht University of Medical Sciences

Full name of responsible person

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

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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

Undecided - It is not yet known if there will be a plan to make this available

Data Dictionary

Undecided - It is not yet known if there will be a plan to make this available

Title and more details about the data/document

Data about primary outcome

When the data will become available and for how long

1 month after results publish

To whom data/document is available

Researchers

Under which criteria data/document could be used

Requested by authenticated scientific centers and universities

From where data/document is obtainable

Dr. Nematollah Ahangar School of Medicine Email: n.ahangar@gums.ac.ir

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

Official request signed by highest executive is mandatory. Moreover, acceptable reasons should be noted

Comments