

# Clinical Trial Protocol

## Iranian Registry of Clinical Trials

09 Jun 2026

### Evaluation of the safety and efficacy of tocilizumab (AryoGen Pharmed Co., Iran) in patients with severe COVID-19

#### Protocol summary

##### Study aim

The evaluation of efficacy and safety of Tocilizumab (AryoGen Pharmed Co.) in patients with severe COVID-19.

##### Design

This is a phase III, open-label, and single-arm study with the sample size of 85 patients.

##### Settings and conduct

This phase-III, single-arm, open-label, multi-center (8 cities of Iran) clinical trial will be conducted in patients with COVID-19 with a follow-up duration of 14 days. Tocilizumab will be administered to the eligible patients on day 1, and the outcomes will be investigated with the 14-day monitoring of the patients.

##### Participants/Inclusion and exclusion criteria

Inclusion criteria: Ability to comprehend and willingness to sign the Informed Consent Form for this study by the patient or his/her guardian; Patients aged  $\geq 18$  years old; Patients with any of the following symptoms: fever (body temperature  $> 37.8^{\circ}\text{C}$ ), shortness of breath, cough, or respiratory rate higher than 30 breaths/min and  $\text{SpO}_2 \leq 93\%$ , with the confirmed diagnosis of SARS-CoV-2 infection based on PCR and/or radiography; Serum Interleukin-6 level  $\geq 3$  times upper limit of normal. Exclusion criteria include hypersensitivity to Tocilizumab or any components of the formulation, hepatic or renal disorders, bone marrow suppression, patients with a high risk of GI perforation, history of tuberculosis, hepatitis B or C, HIV, pregnancy and breastfeeding, and any other active infection.

##### Intervention groups

Intervention group: Subcutaneous (two PFS injections of 162 mg Tocilizumab for patients  $< 100$  kg and three PFS injections of 162 mg Tocilizumab for patients  $> 100$  kg) in addition to the standard treatment. In case of inadequate response, Tocilizumab would be re-administered with a 12-hour interval between injections.

##### Main outcome variables

All-cause mortality rate during the study (date and cause

of death, if applicable)

#### General information

##### Reason for update

To amend the study design (interventions, study endpoints, inclusion criteria, and sample size): based on the state-of-the-art studies in the field of COVID-19 and the fact that improving patient survival is the ultimate goal in these patients, the primary endpoint is changed to the all-cause mortality rate during the study. Secondary endpoints are updated accordingly. Due to the unavailability of the intravenous form, subcutaneous injection was used for all patients. As a result, the interventions section is updated. According to the definition of severe COVID-19 by WHO, respiratory rate higher than 30 breaths/min is added to the inclusion criteria number 3. Based on the significantly elevated IL-6 levels in patients with severe COVID-19 and the mechanism of action of tocilizumab, the inclusion criteria number 4 is changed to serum Interleukin-6 level  $\geq 3$  times upper limit of normal. Based on the first version of the protocol, the trial was going to be conducted in 500 patients with the aim of increasing the patient access to the treatment. Yet, in the current version and due to the increased concern on the safety of tocilizumab treatment in patients with COVID-19, the sample size is calculated to be 85, using Fleming's method (single-arm study) with the statistical power of 90%, alpha level of 5%, and the goal of decreasing the mortality rate by 15% (with the assumption of 60% survival rate with the standard treatment and an increase to 75% by the addition of tocilizumab to the treatment). Moreover, since only severe COVID-19 patients are included in the study, the term "severe" is added to the title. Following the addition of a study center in Semnan and the removal of the centers in Shiraz and Rasht, the total number of cities is changed to 8 in the Summary section.

##### Acronym

##### IRCT registration information

IRCT registration number: **IRCT20150303021315N17**

Registration date: **2020-04-12, 1399/01/24**  
Registration timing: **registered\_while\_recruiting**

Last update: **2020-05-04, 1399/02/15**  
Update count: **1**

**Registration date**  
2020-04-12, 1399/01/24

#### **Registrant information**

##### **Name**

Nassim Anjidani

##### **Name of organization / entity**

Orchid Pharmed

##### **Country**

Iran (Islamic Republic of)

##### **Phone**

+98 21 4347 3000

##### **Email address**

amini@orchidpharmed.com

#### **Recruitment status**

**Recruitment complete**

#### **Funding source**

#### **Expected recruitment start date**

2020-03-15, 1398/12/25

#### **Expected recruitment end date**

2020-07-21, 1399/04/31

#### **Actual recruitment start date**

empty

#### **Actual recruitment end date**

empty

#### **Trial completion date**

empty

#### **Scientific title**

Evaluation of the safety and efficacy of tocilizumab (AryoGen Pharmed Co., Iran) in patients with severe COVID-19

#### **Public title**

Tocilizumab in severe COVID-19

#### **Purpose**

Treatment

#### **Inclusion/Exclusion criteria**

##### **Inclusion criteria:**

Ability to comprehend and willingness to sign the informed consent form for this study by the patient or his/her guardian Patients above 18 years old Patients with fever higher than 37.8 °C, cough, shortness of breath, or respiratory rate higher than 30 breaths/min accompanied by SpO<sub>2</sub> ≤ %93, who have a confirmed diagnosis of SARS-CoV-2 infection using PCR and/or radiography. Serum Interleukin-6 level ≥ 3 times upper limit of normal

##### **Exclusion criteria:**

History of hypersensitivity to Tocilizumab or any components of the formulation Hepatic disease (especially active hepatic diseases and hepatic impairment) Patients with bone marrow suppression (defined as Absolute Neutrophil Count (ANC) below 2000/mm<sup>3</sup> or platelet count below 100,000/mm<sup>3</sup>) Patients with a high risk of gastrointestinal perforation or

with distinct history of GI disorders, e.g., active peptic ulcer and diverticulitis Patients with active or latent tuberculosis Patients with history of active hepatitis B, hepatitis C, HIV, or any known immunodeficiency Pregnant or lactating patients Patients with any other active infection in addition to COVID-19 Patients with any other disease or disorder, which, in the opinion of the investigator will put the subject at risk, if they are enrolled Renal impairment (GFR of below 30ml/min/1.73m<sup>2</sup>)

#### **Age**

From **18 years** old

#### **Gender**

Both

#### **Phase**

3

#### **Groups that have been masked**

*No information*

#### **Sample size**

Target sample size: **85**

#### **Randomization (investigator's opinion)**

Not randomized

#### **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**

National Institution for Medical Research Development

###### **Street address**

No 21, Beesat St., West Fatemi St.

###### **City**

Tehran

###### **Province**

Tehran

###### **Postal code**

1419693111

##### **Approval date**

2020-03-13, 1398/12/23

##### **Ethics committee reference number**

IR.NIMAD.REC.1398.414

### **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**

All-cause mortality rate during the study (date and cause of death, if applicable)

#### **Timepoint**

day 1 through 14

#### **Method of measurement**

Physical Examination

## **Secondary outcomes**

### **1**

#### **Description**

Percent of patients with improvement in oxygenation

#### **Timepoint**

day 1 through 14

#### **Method of measurement**

Percent of patients with at least one step decrease in oxygenation supply; steps: 1. no need for supplemental oxygenation; 2. nasal catheter oxygen inhalation; 3. Mask oxygen inhalation; 4. Noninvasive ventilator oxygen supply; 5. Invasive ventilator oxygen supply.

### **2**

#### **Description**

Duration (days) of hospitalization

#### **Timepoint**

day 1 thorough 14

#### **Method of measurement**

Clinical Evaluation

### **3**

#### **Description**

Changes in body temperature

#### **Timepoint**

Day 1 through 14

#### **Method of measurement**

Physical Examination

### **4**

#### **Description**

The lesions of the pulmonary segment numbers involved in pulmonary CT

#### **Timepoint**

Day 1 and 14

#### **Method of measurement**

CT scan

## **5**

### **Description**

Duration (days) of mechanical ventilation

### **Timepoint**

Day 1 through 14

### **Method of measurement**

Clinical Evaluation

## **6**

### **Description**

Changes in C-reactive protein blood level

### **Timepoint**

Day 1 through 14

### **Method of measurement**

Laboratory Test

## **7**

### **Description**

Changes in Interleukin-6 blood levels

### **Timepoint**

Day 1 through 14

### **Method of measurement**

Laboratory Test

## **8**

### **Description**

Changes in White Blood Cell count

### **Timepoint**

Day 1 through 14

### **Method of measurement**

Laboratory Test

## **9**

### **Description**

Changes in cholesterol level

### **Timepoint**

Day 1 through 14

### **Method of measurement**

Laboratory Test

## **10**

### **Description**

Changes in Aspartate transaminase level

### **Timepoint**

Day 1 through 14

### **Method of measurement**

Laboratory Test

## **11**

### **Description**

Changes in Alanine transaminase level

### **Timepoint**

Day 1 through 14

### **Method of measurement**

Laboratory Test

## 12

### **Description**

Injection site reaction

### **Timepoint**

Day 1 through 14

### **Method of measurement**

Clinical evaluation

## 13

### **Description**

Gastrointestinal perforation

### **Timepoint**

Day 1 through 14

### **Method of measurement**

Clinical evaluation; radiographic imaging in case of clinical symptoms

## 14

### **Description**

Infection

### **Timepoint**

Day 1 through 14

### **Method of measurement**

Clinical evaluation, laboratory test, imaging

## 15

### **Description**

Changes in Platelet level

### **Timepoint**

Day 1 through 14

### **Method of measurement**

Laboratory Test

## 16

### **Description**

Changes in bilirubin level

### **Timepoint**

Day 1 through 14

### **Method of measurement**

Laboratory Test

## 17

### **Description**

Blood pressure

### **Timepoint**

Day 1 through 14

### **Method of measurement**

Blood pressure meter

## **Intervention groups**

### 1

### **Description**

Intervention group: Subcutaneous (two PFS injections of 162 mg Tocilizumab for patients <100 kg and three PFS injections of 162 mg Tocilizumab for patients >100 kg) in

addition to the standard treatment. In case of inadequate response, Tocilizumab would be re-administered with a 12-hour interval between injections.

### **Category**

Treatment - Drugs

## **Recruitment centers**

### 1

#### **Recruitment center**

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

Masih Daneshvari Hospital

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

Payam Tabarsi

##### **Street address**

Darababd St., Shahid Bahonar St., Tehran

##### **City**

Tehran

##### **Province**

Tehran

##### **Postal code**

1956944413

##### **Phone**

+98 21 2610 5050

##### **Email**

payamtabarsi@yahoo.com

##### **Web page address**

<https://nritld.sbmu.ac.ir/index.jsp?fkeyid=&siteid=200&pageid=1419>

### 2

#### **Recruitment center**

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

Imam Reza Hospital

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

Rozita Khodashahi

##### **Street address**

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##### **City**

Mashhad

##### **Province**

Razavi Khorasan

##### **Postal code**

9137913316

##### **Phone**

+98 51 3854 3031

##### **Email**

Rkhodashahi@yahoo.com

##### **Web page address**

<https://emamreza.mums.ac.ir/>

### 3

#### **Recruitment center**

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

Milad Hospital

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

Farzin Khorvash

##### **Street address**

Milad Hospital, Janbazan St, Isfahan.

**City**  
Isfahan  
**Province**  
Isfahan  
**Postal code**  
8174675731  
**Phone**  
+98 31 3777 4001  
**Email**  
khorvash@med.mui.ac.ir  
**Web page address**

#### 4

**Recruitment center**  
**Name of recruitment center**  
Imam Khomeini Hospital  
**Full name of responsible person**  
Roya Ghasemian  
**Street address**  
Valiasr Highway, Joybar three ways  
**City**  
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**Province**  
Mazandaran  
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ict@mazums.ac.ir

#### 5

**Recruitment center**  
**Name of recruitment center**  
Beesat Hospital  
**Full name of responsible person**  
Dr. Behzad Mohsenpour  
**Street address**  
Keshavarz St.,  
**City**  
Sanandaj  
**Province**  
Kurdistan  
**Postal code**  
131651332  
**Phone**  
+98 87 3328 5912  
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Beesat@muk.ac.ir

#### 6

**Recruitment center**  
**Name of recruitment center**  
Imam Khomeini Hospital  
**Full name of responsible person**  
Dr. Mohammad Reza Salehi  
**Street address**  
Dr. Gharib St., Keshavarz Ave.  
**City**  
Tehran

**Province**  
Tehran  
**Postal code**  
1419733141  
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+98 21 6119 0000  
**Email**  
Imamhospital@tums.ac.ir

#### 7

**Recruitment center**  
**Name of recruitment center**  
Shahid Beheshti Hospital  
**Full name of responsible person**  
Dr. Ehsan Sharifipour  
**Street address**  
Shahid Beheshti blvd.  
**City**  
Ghoum  
**Province**  
Ghoum  
**Postal code**  
3719964797  
**Phone**  
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**Email**  
bmc@muq.ac.ir  
**Web page address**  
<https://bmc.muq.ac.ir/index.aspx?siteid=114&fkeyid=&siteid=114&pageid=3554>

#### 8

**Recruitment center**  
**Name of recruitment center**  
Sayyad Shirazi Hospital  
**Full name of responsible person**  
Dr. Nafiseh Abdollahi  
**Street address**  
Shahid Sayyad Shirazi blvd.  
**City**  
Gorgan  
**Province**  
Golestan  
**Postal code**  
4917865086  
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infosayyad@goums.ac.ir

#### 9

**Recruitment center**  
**Name of recruitment center**  
Rasoul Akram Hospital  
**Full name of responsible person**  
Dr. Ali Javad Mousavi  
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Mansoori-Niayesh cross road; Niayesh street;  
Sattarkhan street  
**City**

Tehran  
**Province**  
Tehran  
**Postal code**  
1445613131  
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+98 21 6435 1000  
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info@hrmc.iums.ac.ir

## 10

### Recruitment center

**Name of recruitment center**  
Isabn-e-Maryam Hospital  
**Full name of responsible person**  
Abbas Rezaei  
**Street address**  
Shams Abadi street  
**City**  
Isfahan  
**Province**  
Isfahan  
**Postal code**  
۷۳۴۶۱۸۱۷۴۶  
**Phone**  
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**Web page address**  
<https://isa.mui.ac.ir/>

## 11

### Recruitment center

**Name of recruitment center**  
Kosar Hospital  
**Full name of responsible person**  
Mahboubeh Darban  
**Street address**  
Basij street  
**City**  
Semnan  
**Province**  
Semnan  
**Postal code**  
3519899951  
**Phone**  
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**Email**  
kosarhos@semums.ac.ir  
**Web page address**  
<https://kosarhos.semums.ac.ir/>

## Sponsors / Funding sources

### 1

#### Sponsor

**Name of organization / entity**  
Aryogen Pharmed  
**Full name of responsible person**

Nassim Anjidani  
**Street address**  
Next to Tajbakhsh street, 24th Kilometer of Makhsous road  
**City**  
Garm Darreh  
**Province**  
Alborz  
**Postal code**  
56145226  
**Phone**  
+98 26 3610 6480  
**Email**  
contact@aryogen.com

**Grant name**  
**Grant code / Reference number**  
**Is the source of funding the same sponsor organization/entity?**  
Yes  
**Title of funding source**  
Aryogen Pharmed  
**Proportion provided by this source**  
100  
**Public or private sector**  
Private  
**Domestic or foreign origin**  
Domestic  
**Category of foreign source of funding**  
*empty*  
**Country of origin**  
**Type of organization providing the funding**  
Industry

## Person responsible for general inquiries

### Contact

**Name of organization / entity**  
OrchidPharmed Co.  
**Full name of responsible person**  
Nassim Anjidani  
**Position**  
Medical Department Manager  
**Latest degree**  
Medical doctor  
**Other areas of specialty/work**  
Medical Pharmacy  
**Street address**  
No 42, Attar street, Vanak square  
**City**  
Tehran  
**Province**  
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**Postal code**  
1651655  
**Phone**  
009843473000  
**Email**  
anjidani.n@orchipharmed.com

## Person responsible for scientific

## inquiries

### Contact

**Name of organization / entity**

Shahid Beheshti University of Medical Sciences

**Full name of responsible person**

Dr. Payam Tabarsi

**Position**

Professor

**Latest degree**

Ph.D.

**Other areas of specialty/work**

Infectious diseases

**Street address**

Masih Daneshvari Hospital, Darabad St., Bahonar St.,

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**Email**

payamtabarsi@yahoo.com

## Person responsible for updating data

### Contact

**Name of organization / entity**

Orchid Pharmed

**Full name of responsible person**

Nassim Anjidani

**Position**

Medical Manager

**Latest degree**

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**Other areas of specialty/work**

Medical Pharmacy

**Street address**

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+98 21 8856 2862

**Email**

anjidani.n@orchidpharmed.com

## Sharing plan

**Deidentified Individual Participant Data Set (IPD)**

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

**Study Protocol**

Undecided - It is not yet known if there will be 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**

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

**Clinical Study Report**

Undecided - It is not yet known if there will be 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