

# Trial to study the effect of Curvic® in vaccinated population for COVID-19

<b>Submission date</b> 14/03/2022	<b>Recruitment status</b> No longer recruiting	<input type="checkbox"/> Prospectively registered <input checked="" type="checkbox"/> Protocol
<b>Registration date</b> 19/04/2022	<b>Overall study status</b> Completed	<input type="checkbox"/> Statistical analysis plan <input type="checkbox"/> Results
<b>Last Edited</b> 20/04/2022	<b>Condition category</b> Infections and Infestations	<input type="checkbox"/> Individual participant data <input type="checkbox"/> Record updated in last year

## Plain English summary of protocol

### Background and study aims

Vaccination is protective against COVID-19 infections. Based on recent studies by Shreepad Shree Vallabh SSV Phytopharmaceuticals, it has been scientifically proven that Curvic® is effective against all species of coronavirus by blocking the Main Protein (Mpro). The efficacy of Curvic® has been proven in quarantined patients as an immunity booster and also in a randomized double-blind trial for the management of SARS-COV-2 (COVID-19) infection. We plan to conduct the current study to evaluate the role of Curvic® in the COVID-19 vaccinated population.

### Who can participate?

Adult male or female volunteers of age 18-65 years (inclusive), who are eligible for vaccination for COVID-19

### What does the study involve?

After vaccination, participants will be randomly allocated to receive Curvic® or not. Blood samples will be taken at baseline and 120 days

### What are the possible benefits and risks of participating?

The supplement Curvic® may benefit participants who are vaccinated for COVID-19. Curvic®, being a nutraceutical supplement, possesses no risk to the participants consuming it. This has been proven in various clinical trials carried out previously for the formulation.

### Where is the study run from?

Sukhkarta Hospital (India)

### When is the study starting and how long is it expected to run for?

March 2021 to February 2022

### Who is funding the study?

Shreepad Shree Vallabh SSV Phytopharmaceuticals (India)

Who is the main contact?

Dr Yogesh Dound, yogesh\_dound@yahoo.com

## Contact information

### Type(s)

Scientific

### Contact name

Dr Yogesh Dound

### ORCID ID

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### Contact details

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## Additional identifiers

### EudraCT/CTIS number

Nil known

### IRAS number

### ClinicalTrials.gov number

Nil known

### Secondary identifying numbers

SSV-SF-01/2021

## Study information

### Scientific Title

A randomized controlled trial to study the immunogenicity and safety of Curvic® in a vaccinated population for COVID-19

### Acronym

SSV-SF-01/2021

### Study objectives

Vaccination is protective against COVID 19 infections. Based on recent studies by Shreepad Shree Vallabh SSV Phytopharmaceuticals, it has been scientifically proven that Curvic® is effective against all species of coronavirus by blocking the Main Protein (Mpro). The efficacy of Curvic® has been proven in quarantined patients as an immunity booster and also in a randomized double-blind trial for the management of SARS-COV-2 (COVID-19) infection. We plan to conduct the current study to evaluate the role of Curvic® in the vaccinated population for COVID 19.

### **Ethics approval required**

Old ethics approval format

### **Ethics approval(s)**

Approved 21/03/2021, Navsanjeevani Hospital Ethics Committee (Nashik, Maharashtra, India; +91 9822574157; navsanjeevaniec@gmail.com), ref: none provided

### **Study design**

Interventional randomized controlled trial

### **Primary study design**

Interventional

### **Secondary study design**

Randomised controlled trial

### **Study setting(s)**

Hospital

### **Study type(s)**

Treatment

### **Participant information sheet**

See additional files

### **Health condition(s) or problem(s) studied**

Vaccinated for COVID-19

### **Interventions**

The study shall include 50 healthy volunteers eligible for vaccination for COVID-19. These volunteers will be screened for Absolute Lymphocyte Count (CD3+, CD4+ and CD8+) and Anti SARS-CoV-2 spike protein (S1 /S2) IgG. Once vaccinated, they will be divided in two groups:

Group A: (N=25) Curvic® given to vaccinated population for COVID-19 for 120 days

Group B: (N=25) Only vaccinated population for COVID-19

Once vaccinated they will follow the protocol for post vaccination. Based on the type of vaccination, whichever is the due date for 2nd vaccination will be followed by the respective volunteer. After completion of 120 days from first dose of vaccination, the volunteers will be again screened for Absolute Lymphocyte Count (CD3+, CD4+ and CD8+) and Anti SARS-CoV-2 spike protein (S1 /S2) IgG in both the groups.

### **SUBJECT RECRUITMENT:**

Healthy volunteers will be recruited into the study based on the inclusion criteria and the data shall be collected i.e. Demographics, comorbidities, date of vaccination will be entered in excel.

They will be closely followed up to 120 days from the first dose of vaccination for COVID-19 to observe for any adverse reactions.

**RANDOMIZATION:**

Computer generated randomization

**DISCONTINUATION CRITERIA:**

People who become infected during the study period and lost to follow up are excluded.

**Intervention Type**

Supplement

**Primary outcome measure**

Measured by blood test at baseline and 120 days:

1. Absolute lymphocyte count (CD3+, CD4+ and CD8+)
2. Anti-SARS-CoV-2 spike protein (S1 /S2) IgG

**Secondary outcome measures**

Safety of Curvic® with or without vaccine assessed through clinical biochemistry and by symptomatology at baseline and 120 days

**Overall study start date**

21/03/2021

**Completion date**

28/02/2022

## **Eligibility**

**Key inclusion criteria**

1. Adult male or female human volunteer of age 18-65 years (inclusive)
2. Eligible for vaccination for COVID-19
3. Willing and able to provide written, signed and dated informed consent
4. Had no medical history or evidence of COVID-2019
5. Had no acute infections and/or respiratory diseases within 14 days before enrollment.
6. Had no evidence of vaccine-induced reactions or complications after receiving immunobiological products in the past medical history.
7. Willing to give consent to use effective contraception methods during the study
8. Have a negative urine pregnancy test at the screening visit (for child-bearing aged women)
9. Have negative human immunodeficiency virus (HIV 1 & 2), syphilis, hepatitis B and C test results

**Participant type(s)**

Healthy volunteer

**Age group**

Adult

**Lower age limit**

18 Years

**Upper age limit**

65 Years

**Sex**

Both

**Target number of participants**

50

**Total final enrolment**

50

**Key exclusion criteria**

1. Aged <18 years and >65 years
2. Any vaccination/immunization within 30 days before the enrollment
3. Any treatment with steroids (except hormonal contraceptives) and/or immunoglobulins or other blood products within 30 days before the enrollment
4. Any treatment with immunosuppressive therapy within 3 months before the enrollment
5. Any drug allergy (anaphylactic shock, Quincke's edema, polymorphic exudative eczema, atopy, serum disease), hypersensitivity or allergic reaction to immunobiological products, known allergic reactions to study drug components, acute exacerbation of allergic diseases on enrollment day
6. Any neoplasms in the past medical history
7. Donated blood or plasma within 3 months before the enrollment
8. Any history or evidence of splenectomy
9. Any immunodeficiency state
10. Any history or evidence of anorexia or protein deficiency of any origin
11. Alcohol or drug addiction in the past medical history
12. Participation in any other interventional clinical trial within 3 months
13. Any other condition that the study physician considers as a barrier to the trial completion as per the protocol
14. Pregnancy or breast-feeding
15. Subjects who are tested positive for Coronavirus disease through RT-PCR SARS CoV-2 Test and positive anti-COVID antibodies within 3 months before the enrollment

**Date of first enrolment**

01/04/2021

**Date of final enrolment**

25/12/2021

**Locations****Countries of recruitment**

India

**Study participating centre**

Sukhkarta Hospital

1st floor, Bhagchand Complex

Above Abhishek Agro  
Dindori Naka  
Panchavati  
Nashik  
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## Sponsor information

### Organisation

Shreepad Shree Vallabh SSV Phytopharmaceuticals

### Sponsor details

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director.ssvphytopharma@gmail.com

### Sponsor type

Industry

### Website

<http://www.ssvphytopharma.com>

## Funder(s)

### Funder type

Industry

### Funder Name

Shreepad Shree Vallabh SSV Phytopharmaceuticals

## Results and Publications

### Publication and dissemination plan

Planned publication in a high-impact peer-reviewed journal

## Intention to publish date

05/03/2022

## Individual participant data (IPD) sharing plan

Raw data can be shared upon request by contacting [yogesh\\_dound@yahoo.com](mailto:yogesh_dound@yahoo.com)

## IPD sharing plan summary

Available on request

## Study outputs

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
<a href="#">Participant information sheet</a>	English version 1	09/03/2021	16/03/2022	No	Yes
<a href="#">Participant information sheet</a>	Marathi version 1	09/03/2021	16/03/2022	No	Yes
<a href="#">Protocol file</a>	version 1	09/03/2021	16/03/2022	No	No