



Clinical Trial Details (PDF Generation Date :- Wed, 07 Jun 2023 17:54:38 GMT)

CTRI Number	CTRI/2020/08/027170 [Registered on: 15/08/2020] - Trial Registered Prospectively	
Last Modified On	12/06/2021	
Post Graduate Thesis	No	
Type of Trial	Interventional	
Type of Study	Vaccine	
Study Design	Randomized, Parallel Group Trial	
Public Title of Study	Study to check the safety and immune response of a COVID-19 vaccine in healthy Indian adults.	
Scientific Title of Study	A Phase 2/3, Observer-Blind, Randomized, Controlled Study to Determine the Safety and Immunogenicity of Covishield (COVID-19 Vaccine) in Healthy Indian Adults	
Secondary IDs if Any	Secondary ID	Identifier
	ICMR/SII-COVISHIELD Version 4.0 dated 14 Oct 2020	Protocol Number
Details of Principal Investigator or overall Trial Coordinator (multi-center study)	Details of Principal Investigator	
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	Designation	
	Affiliation	
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Source of Monetary or Material Support	Source of Monetary or Material Support			
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	> Serum Institute of India Private Limited 212/2, Hadapsar, Pune – 411 028, India			
Primary Sponsor	Primary Sponsor Details			
	Name	Serum Institute of India Private Limited		
	Address	212/2, Off Soli Poonawalla Road, Hadapsar, Pune – 411 028, India		
	Type of Sponsor	Pharmaceutical industry-Indian		
Details of Secondary Sponsor	Name	Address		
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Countries of Recruitment	List of Countries			
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Details of Ethics Committee

Name of Committee	Approval Status	Date of Approval	Is Independent Ethics Committee?
Mahatma Gandhi Institute of Medical Sciences, Institutional Ethics Committee, Sewagram	Approved	26/10/2020	No
Ethics Committee Jehangir Clinical Development Center Pvt.Ltd, Pune	Approved	22/10/2020	No
Ethics Committee Rajendra Memorial Research Institute of Medical Sciences,	Approved	29/10/2020	No



Patna			
IEC King George Hospital, Visakhapatnam	Approved	28/10/2020	No
Institutional Ethics Committee - TNGMSSH, Chennai	Approved	13/08/2020	No
Institutional Ethics Committee of B. J. Government Medical College and Sassoon General Hospital, Pune	Approved	26/10/2020	No
Institutional Ethics Committee Seth GS Medical College and KEM Hospital, Mumbai.	Approved	28/10/2020	No
Institutional Ethics Committee Sri Ramachandra Institute of Higher Education and Research, Chennai	Approved	29/10/2020	No
Institutional Ethics Committee, BVDU, Pune	Approved	14/08/2020	No
Institutional Ethics Committee, Government Medical College, Nagpur	Approved	26/10/2020	No
Institutional Ethics Committee, JSS Medical College, Mysore	Approved	24/10/2020	No
Institutional Ethics Committee, PGIMER, Chandigarh	Approved	29/10/2020	No
Institutional Ethics Committee, T N Medical College & BYL Nair Hospital, Mumbai	Approved	27/10/2020	No
KEM Hospital Research Centre Ethics Committee, Pune	Approved	24/10/2020	No

Regulatory Clearance Status from DCGI

Status	Date
Approved/Obtained	16/10/2020

Health Condition / Problems Studied

Health Type	Condition
Healthy Human Volunteers	Prevention of COVID-19 infection

Intervention / Comparator Agent

Type	Name	Details
Intervention	Covishield (SII-ChAdOx1 nCoV-19)	Covishield will be administered as 2 dose schedule on Days 1 and 29 as 0.5 ml dose intramuscularly.
Comparator Agent	Oxford/AZ-ChAdOx1 nCoV-19 vaccine	Oxford/AZ-ChAdOx1 nCoV-19 vaccine will be administered as 2 dose schedule on Days 1 and 29 as 0.5 ml dose



Comparator Agent	Placebo	intramuscularly. Placebo will be administered as 2 dose schedule on Days 1 and 29 as 0.5 ml dose intramuscularly.
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Inclusion Criteria

Inclusion Criteria	
Age From	18.00 Year(s)
Age To	99.00 Year(s)
Gender	Both
Details	1. Healthy adults aged more than or equal to 18 years of either sex. 2. Written informed consent by participants. 3. The participant is resident of the study area and is willing to comply with study protocol requirements. 4. Healthy, as determined by medical history and physical examination. 5. Female participants of childbearing potential must have a negative urine pregnancy test within 24 hours prior to study vaccine administration.

Exclusion Criteria

Exclusion Criteria	
Details	1. Acute illness with or without fever at the time of study vaccine administration 2. History of laboratory confirmed COVID-19 disease in household contact or close workplace contact 3. IgG seropositivity to SARS-CoV-2 4. History or currently positive for SARS-CoV-2 by RT-PCR 5. History of severe allergic reactions after previous vaccinations or hypersensitivity to any component of study vaccines 6. Any confirmed or suspected condition with impaired/altered function of immune system

Method of Generating Random Sequence

Computer generated randomization

Method of Concealment

Centralized

Blinding/Masking

Participant, Investigator, Outcome Assessor and Data-entry Operator Blinded

Primary Outcome

Outcome	Timepoints
1. Occurrence of causally related SAEs throughout the study duration following vaccination 2. Ratio of GMTs of anti-S IgG antibodies	1. Throughout the study duration following vaccination 2. 28 days after the second vaccination

Secondary Outcome

Outcome	Timepoints
Occurrence of solicited local and/or systemic adverse events (AEs)	7 days following each vaccination
Occurrence of unsolicited adverse events	28 days following each vaccination
Occurrence of serious adverse events (SAEs)	Throughout the study duration following vaccination

Target Sample Size

Total Sample Size=1600
Sample Size from India=1600
Final Enrollment numbers achieved (Total)=0
Final Enrollment numbers achieved (India)=1600

Phase of Trial

Phase 2/ Phase 3

Date of First Enrollment (India)

24/08/2020

Date of First Enrollment (Global)

No Date Specified

Estimated Duration of

Years=0



Trial	Months=7 Days=0
Recruitment Status of Trial (Global)	Not Applicable
Recruitment Status of Trial (India)	Completed
Publication Details	Nil
Brief Summary	<p>This is a Phase 2/3, observer-blind, randomised, controlled study in healthy adults in India, for comparison of the safety of COVISHIELD with Oxford/AZ-ChAdOx1 nCoV-19 and Placebo, and immunogenicity with Oxford/AZ-ChAdOx1 nCoV-19 in prevention of SARS CoV-2 infection. A total of 1600 eligible participants of more than or equal to 18 years of age will be enrolled the study. Of these 400 participants will be part of immunogenicity cohort and will be randomly assigned in a 3:1 ratio to receive either COVISHIELD or Oxford/AZ-ChAdOx1 nCoV-19, respectively. The remaining 1200 participants from safety cohort will be randomly assigned in a 3:1 ratio to receive either COVISHIELD or Placebo, respectively.</p>